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(57) Abstract

The invention relates to recombinant polypeptides and peptides and particularly to the polypeptide containing in its polypeptidic chain the following amino acid sequence: the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (194) represented in fig. 4a and fig. 4b. The polypeptides and peptides of the invention can be used for the diagnostic of tuberculosis, and can also be part of the active principle in the preparation of vaccine against tuberculosis.

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RECOMBINANT POLYPEPTIDES AND PEPTIDES, NUCLEIC ACIDS CODING FOR THE SAME AND USE OF THESE POLYPEPTIDES AND PEPTIDES IN THE DIAGNOSTIC OF TUBERCULOSIS

The invention relates to recombinant polypeptides and peptides, which can be used for the diagnosis of tuberculosis. The invention also relates to a process for preparing the above-said polypeptides and peptides, which are in a state of biological purity such that they can be used as part of the active principle in the preparation of vaccines against tuberculosis.

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It also relates to nucleic acids coding for said polypeptides and peptides.

Furthermore, the invention relates to the <u>in vitro</u> diagnostic methods and kits using the above-said polypeptides and peptides and to the vaccines containing the above-said polypeptides and peptides as active principle against tuberculosis.

By "recombinant polypeptides or peptides" it is to be understood that it relates to any molecule having a polypeptidic chain liable to be produced by genetic engineering, through transcription and translation, of a corresponding DNA sequence under the control of appropriate regulation elements within an efficient cellular host. Consequently, the expression "recombinant polypeptides" such as is used herein does not exclude the possibility for the polypeptides to comprise other groups, such as glycosylated groups.

The term "recombinant" indeed involves the fact that the polypeptide has been produced by genetic engineering, particularly because it results from the expression in a cellular host of the corresponding nucleic acid sequences which have previously been introduced into the expression vector used in said host.

Nevertheless, it must be understood that this expression does not exclude the possibility for the polypeptide to be produced by a different process, for instance by classical chemical synthesis according to methods used in the protein synthesis or by proteolytic cleavage of larger molecules.

The expression "biologically pure" or "biological purity" means on the one hand a grade of purity such that the recombinant polypeptide can be used for the production of vaccinating compositions and on the other hand the absence of contaminants, more particularly of natural contaminants.

Tuberculosis remains a major disease in developing countries. The situation is dramatic in some countries, particularly where high incidence of tuberculosis among AIDS patients represents a new source of dissemination of the disease.

Tuberculosis is a chronic infectious disease in which cell-mediated immune mechanisms play an essential role both for protection against and control of the disease.

Despite BCG vaccination, and some effective drugs, tuberculosis remains a major global problem. Skin testing with tuberculin PPD (protein-purified derivative) largely used for screening of the disease is poorly specific, due to cross reactivity with other pathogenic or environmental saprophytic mycobacteria.

Moreover, tuberculin PPD when used in serological tests (ELISA) does not allow to discriminate between patients who have been vaccinated by BCG, or those who have been primo-infected, from those who are developing evolutive tuberculosis and for whom an early and rapid diagnosis would be necessary.

A protein with a molecular weight of 32-kDa has been purified (9) from zinc deficient Mycobacterium bovis BCG culture filtrate (8). This 32-kDa protein of

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 \underline{M} . bovis BCG has been purified from Sauton zinc deficient culture filtrate of \underline{M} . bovis BCG using successively hydrophobic chromatography on Phenyl-Sepharose, ion exchange on DEAE-Sephacel and molecular sieving on Sephadex G-100. The final preparation has been found to be homogeneous as based on several analyses. This P_{32} protein is a constituent of BCG cells grown in normal conditions. It represents about 3% of the soluble fraction of a cellular extract, and appears as the major protein released in normal Sauton culture filtrate. This protein has been found to have a molecular weight of 32 000 by SDS-polyacrylamide gel electrophoresis and by molecular sieving.

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The $\mathrm{NH_2}\text{-terminal}$ amino acid sequence of the 32-kDa protein of M. bovis BCG (Phe-Ser-Arg-Pro-Gly-Leu) is identical to that reported for the MPB 59 protein purified from M. bovis BCG substrain Tokyo (34).

Purified P_{32} of <u>M. bovis</u> BCG has been tested by various cross immunoelectrophoresis techniques, and has been shown to belong to the antigen 85 complex in the reference system for BCG antigens. It has been more precisely identified as antigen 85A in the closs reference system for BCG antigens (7).

Increased levels of immunoglobulin G antibodies towards the 32-kDa protein of <u>M. bovis</u> BCG could be detected in 70% of tuberculous patients (30).

Furthermore, the 32-kDa protein of \underline{M} . bovis BCG induces specific lymphoproliferation and interferon-(IFN- γ) production in peripheral blood leucocytes from patients with active tuberculosis (12) and PPD-positive healthy subjects. Recent findings indicate that the amount of 32-kDa protein of \underline{M} . bovis BCG-induced IFN- γ in BCG-sensitized mouse spleen cells is under probable H-2 control (13). Finally, the high affinity of mycobacteria for fibronectin is related to proteins of the BCG 85 antigen complex (1).

Matsuo et al. (17) recently cloned the gene encoding the antigen α , a major protein secreted by BCG (substrain Tokyo) and highly homologous to MPB 59 antigen in its NH₂-terminal amino acid sequence, and even identical for its first 6 amino acids: Phe-Ser-Arg-Pro-Gly-Leu.

This gene was cloned by using a nucleotide probe homologous to the N-terminal amino acid sequence of antigen α , purified from <u>M. tuberculosis</u> as described in Tasaka, H. et al., 1983. "Purification and antigenic specificity of alpha protein (Yoneda and Fukui) from Mycobacterium tuberculosis and Mycobacterium intracellulare. Hiroshima J. Med. Sci. <u>32</u>, 1-8.

The presence of antigens of around 30-32-kDa, named antigen 85 complex, has been revealed from electrophoretic patterns of proteins originating from culture media of mycobacteria, such as Mycobacterium tuberculosis. By immunoblotting techniques, it has been shown that these antigens cross-react with rabbit sera raised against the 32-kDa protein of BCG (8).

A recent study reported on the preferential humoral response to a 30-kDa and 31-kDa antigen in lepromatous leprosy patients, and to a 32-kDa antigen in tuberculoid leprosy patients (24).

It has also been found that fibronectin (FN)-binding antigens are prominent components of short-term culture supernatants of Mycobacterium tuberculosis. In 3-day-old supernatants, a 30-kilodalton (kDa) protein was identified as the major (FN)-binding molecule. In 21-day-old supernatants, FN was bound to a double protein band of around 30 to 32-kDa, as well as to a group of antigens of larger molecular mass (57 to 60 kDa)(1).

In other experiments, recombinant plasmids containing DNA from Mycobacterium tuberculosis were transformed into Escherichia coli, and three colonies

were selected by their reactivity with polyclonal antisera to <u>M. tuberculosis</u>. Each recombinant produced 35- and 53-kilodalton proteins (35K and 53K proteins, respectively) ("Expression of Proteins of Mycobacterium tuberculosis in Escherichia coli and Potential of Recombinant Genes and Proteins for Development of Diagnostic Reagents", Mitchell L Cohen et al., Journal of Clinical Microbiology, July 1987, p.1176-1180).

Concerning the various results known to date, the physico-chemical characteristics of the antigen P_{32} of Mycobacterium tuberculosis are not precise and, furthermore, insufficient to enable its unambiguous identifiability, as well as the characterization of its structural and functional elements.

Moreover, the pathogenicity and the potentially infectious property of <u>M. tuberculosis</u> has hampered research enabling to identify, purify and characterize the constituents as well as the secretion products of this bacteria.

An aspect of the invention is to provide recombinant polypeptides which can be used as purified antigens for the detection and control of tuberculosis.

Another aspect of the invention is to provide nucleic acids coding for the peptidic chains of biologically pure recombinant polypeptides which enable their preparation on a large scale.

Another aspect of the invention is to provide antigens which can be used in serological tests as an in vitro rapid diagnostic of tuberculosis.

Another aspect of the invention is to provide a rapid <u>in vitro</u> diagnostic means for tuberculosis, enabling it to discriminate between patients suffering from an evolutive tuberculosis from those who have been vaccinated against BCG or who have been primo-infected.

Another aspect of the invention is to provide nucleic probes which can be used as <u>in vitro</u> diagnostic

reagent for tuberculosis, as well as <u>in vitro</u> diagnostic reagent for identifying <u>M. tuberculosis</u> from other strains of mycobacteria.

The recombinant polypeptides of the invention contain in their polypeptidic chain one at least of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (275) to the extremity

constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,

and the peptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids in so far as this modification does not alter the following properties:

the polypeptides react with rabbit polyclonal antiserum raised against the protein of 32-kDa of $\underline{\text{M.}}$ bovis BCG culture filtrate, and/or

react selectively with human sera from tuberculosis patients and particularly patients developing an evolutive tuberculosis at an early stage,

and/or react with the amino acid sequence extending from the extremity constituted by amino acid at position (1), to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b.

On figures 3a and 3b:

- X represents G or GG,
- Y represents C or CC,
- Z represents C or G,
- W represents C or G and is different from Z,
- K represents C or CG,
- L represents G or CC,
- a₁-b₁ represents ALA-ARG or GLY-ALA-ALA,
- a2 represents arg or gly,
- $-a_3-b_3-c_3-d_3-e_3-f_3-$ represents

his-trp-val-pro-arg-pro or

ala-leu-gly-ala,

- a represents pro or pro-asn-thr,
- a₅ represents pro or ala-pro.

The recombinant polypeptides of the invention contain in their polypeptidic chain one at least of the following amino acid sequences:

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- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (275) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- and the peptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids in so far as

this modification does not alter the following properties:

the polypeptides react with rabbit polyclonal antiserum raised against the protein of 32-kDa of $\underline{\text{M.}}$ bovis BCG culture filtrate, and/or

react selectively with human sera from tuberculosis patients and particularly patients developing an evolutive tuberculosis at an early stage,

and/or react with the amino acid sequence extending from the extremity constituted by amino acid at position (1), to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b.

The recombinant polypeptides of the invention contain in their polypeptidic chain one at least of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-30) to the extremity constituted by amino acid at position (-1) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 5, or

- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 5, or

- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 5, or

- the one extending from the extremity constituted by amino acid at position (275) to the extremity constituted by amino acid at position (295) represented on fig. 5,

and the peptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids in so far as this modification does not alter the following properties:

the polypeptides react with rabbit polyclonal antiserum raised against the protein of 32-kDa of <u>M. bovis</u> BCG culture filtrate, and/or

react selectively with human sera from tuberculosis patients and particularly patients developing an evolutive tuberculosis at an early stage,

and/or react with the amino acid sequence extending from the extremity constituted by amino acid at position (1), to the extremity constituted by amino acid at position (295) represented on fig. 5.

Advantageous polypeptides of the invention are characterized by the fact that they react with rabbit polyclonal antiserum raised against the protein of 32-kDa of M. bovis BCG culture filtrate, hereafter designated by "P₃₂ protein of BCG".

Advantageous polypeptides of the invention are characterized by the fact that they selectively react with human sera from tuberculous patients and

particularly patients developing an evolutive tuberculosis at an early stage.

Hereafter is given, in a non limitative way a for preparing rabbit polyclonal antiserum raised against the P_{32} protein of BCG and a test for reaction between the of the evidence giving invention and said polypeptides of the polyclonal antiserum raised against the P32 protein of BCG.

1) process for preparing rabbit polyclonal antiserum raised against the P_{32} protein of BCG:

Purified P_{32} protein of BCG from culture filtrate is used.

a) Purification of protein P_{32} of BCG: P_{32} protein can be purified as follows:

The bacterial strains used are <u>M. bovis</u> BCG substrains 1173P2 (Pasteur Institute, Paris) and GL2 (Pasteur Institute, Brussels).

The culture of bacteria is obtained as follows:

Mycobacterium bovis BCG is grown as a pellicle on Sauton medium, at 37.5°C for 14 days. As the medium is prepared with distilled water, zinc sulfate is added to the final concentration of 5 μ M (normal Sauton medium) (De Bruyn J., Weckx M., Beumer-Jochmans M.-P. Effect of zinc deficiency on Mycobacterium tuberculosis var. bovis (BCG). J. Gen. Microbiol. 1981; 124:353-7). When zinc deficient medium was needed, zinc sulfate is omitted.

The filtrates from zinc deficient cultures are obtained as follows:

The culture medium is clarified by decantation. The remaining bacteria are removed by filtration through Millipak 100 filter unit (Millipore Corp., Bedford, Mass.). When used for purification, the filtrate is adjusted to 20 mM in phosphate, 450 mM in NaCl, 1 mM in EDTA, and the pH is brought to 7.3 with

5 M HCl before sterile filtration.

out by analysis is carried protein polyacrylamide gel electrophoresis. dodecyl Sodium sulfate polyacrylamide gel electrophoresis (SDS-PAGE) was done on 13% (w/v) acrylamide-containing gels as described by Laemmli UK. (Cleavage of structural head during the assembly of the proteins bacteriophage T4. Nature 1970; 227:680-5). The gels are stained with Coomassie Brilliant Blue R-250 and for quantitative analysis, scanned at 595 nm with a DU8 Beckman spectrophotometer. For control of purity the gel is revealed with silver stain (Biorad Laboratories, Richmond, Calif.).

The purification step of P_{32} is carried out as follows:

Except for hydrophobic chromatography on Phenyl-Sepharose, all buffers contain Tween 80 (0.005% final concentration). The pH is adjusted to 7.3 before sterilization. All purification steps are carried out at +4°C. Elutions are followed by recording the absorbance at 280 nm. The fractions containing proteins are analysed by SDS-PAGE.

- (i) The treated filtrate from a 4 liters zincdeficient culture, usually containing 125 to 150 mg protein per liter, is applied to a column (5.0 by 5.0 Phenyl-Sepharose CL-4B (Pharmacia which is previously Sweden), Chemicals, Uppsala, with phosphate buffer 20 mM equilibrated containing 0.45 M NaCl and 1 mM EDTA, at a flow rate of 800 ml per hour. The gel is then washed with one column volume of the same buffer to remove unfixed material and successively with 300 ml of 20 mM and 4 mM PB and 10% ethanol (v/v). The P_{32} appears in the fraction eluted with 10% ethanol.
- (ii) After the phosphate concentration of this fraction has been brought to 4 mM, it is applied to a column (2.6 by 10 cm) of DEAE-Sephacel (Pharmacia Fine

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Chemicals), which is equilibrated with 4 mM PB. After washing with the equilibrating buffer the sample is eluted with 25 mM phosphate at a flow rate of 50 ml per hour. The eluate is concentrated in a 202 Amicon stirred cell equipped with a PM 10 membrane (Amicon Corp., Lexington, Mass.).

- (iii) The concentrated material is submitted to 4 mg of P_{32} protein of BCG (soluble extract) or molecular sieving on a Sephadex G-100 (Pharmacia) column (2.6 by 45 cm) equilibrated with 50 mM PB, at a flow rate of 12 ml per hour. The fractions of the peak giving one band SDS-PAGE are pooled. The purity of the final preparation obtained is controlled by SDS-PAGE followed by silverstaining and by molecular sieving by (Pharmacia) column (12.0)Superose 12 equilibrated with 50 mM PB containing 0.005% Tween 80 at a flow rate of 0.2 ml/min. in the Fast Protein Liquid Chromatography system (Pharmacia). Elution is followed by recording the absorbance at 280 nm and 214 nm.
- b) Preparation of rabbit polyclonal antiserum raised against the P_{32} protein of BCG :

400 μ g of purified P₃₂ protein of BCG per ml physiological saline are mixed with one volume of incomplete Freund's adjuvant. The material is homogenized and injected intradermally in 50 μ l doses delivered at 10 sites in the back of the rabbits, at 0, 4, 7 and 8 weeks (adjuvant is replaced by the diluent for the last injection). One week later, the rabbits are bled and the sera tested for antibody level before being distributed in aliquots and stored at -80°C;

2) test for giving evidence of the reaction between the polypeptides of the invention and said rabbit polyclonal antiserum raised against the P_{32} protein of BCG:

the test used was an ELISA test; the ELISA for antibody determination is based on the method of Engvall and Perlmann (Engvall, E., and P. Perlmann. 1971. Enzyme-linked immunosorbent assay (ELISA). Quantitative assay of immunoglobulin G. Immunochemistry 8:871-874)

Immulon Microelisa plates (Dynatech, Kloten, Switzerland) are coated by adding to each well 1 μ g of one of the polypeptides of the invention in 100 µl Tris hydrochloride buffer 50 mM (pH 8.2). After incubation for 2 h at 27°C in a moist chamber, the plates are kept overnight at 4°C. They are washed four times with 0.01 M phosphate-buffered saline (pH 7.2) containing 0.05% Tween 20 by using a Titertek microplate washer (Flow Laboratories. Brussels. Belgium). Blocking is done with 0.5% gelatin in 0.06 M carbonate buffer (pH 9.6) for 1 h. Wells are then washed as before, and 100 μ l of above mentioned serum diluted in phosphatebuffered saline containing 0.05% Tween 20 and 0.5% gelatin is added. According to the results obtained in preliminary experiments, the working dilutions are set at 1:200 for IqG, 1:20 for IqA and 1:80 for IqM determinations. Each dilution is run in duplicate. After 2 h of incubation and after the wells are washed, they are filled with 100 μ l of peroxidase-conjugated rabbit immunoglobulins directed against human IgG, IgA or IgM (Dakopatts, Copenhagen, Denmark), diluted 1:400, 1:400 and 1:1.200, respectively in phosphate-buffered saline containing 0.05% Tween 20 and 0.5% gelatin and incubated for

90 min. After the wash, the amount of peroxidase bound to the wells is quantified by using a freshly prepared solution of o-phenylenediamine (10 mg/100 ml) and hydrogen peroxide (8 μ l of 30% H₂O₂ per 100 ml) in 0.15 M citrate buffer (pH 5.0) as a substrate. The enzymatic reaction is stopped with 8 N H₂SO₄ after

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15 min. of incubation. The optical density is read at 492 nm with a Titertek Multiskan photometer (Flow Laboratories).

Wells without sera are used as controls for the conjugates. Each experiment is done by including on each plate one negative and two positive reference sera with medium and low antibody levels to correct for plate-to-plate and day-to-day variations. The antibody concentrations are expressed as the optical density values obtained after correction of the readings according to the mean variations of the reference sera.

Hereafter is also given in a non limitative way, a test for giving evidence of the fact that polypeptides of the invention are recognized selectively by human sera from tuberculous patients.

This test is an immunoblotting (Western blotting) analysis, in the case where the polypeptides of the invention are obtained by recombinant techniques. This test can also be used for polypeptides of the invention obtained by a different preparation process. sulfate-polyacrylamide gel dodecyl electrophoresis, polypeptides of the invention blotted onto nitrocellulose membranes (Hybond (Amersham)) as described by Towbin et al. (29). The expression of polypeptides of the invention fused to β -galactosidase in E. coli Y1089, is visualized by the binding of a polyclonal rabbit anti-32-kDa BCG protein serum (1:1,000) or by using a monoclonal anti- β -The secondary (Promega). galactosidase antibody anti-rabbit (alkaline phosphatase antibody immunoglobulin G and anti-mouse alkaline phosphatase immunoglobulin G conjugates, respectively) is diluted as recommended by the supplier (Promega).

In order to identify selective recognition of polypeptides of the invention and of fusion proteins of the invention by human tuberculous sera, nitrocellulose

sheets are incubated overnight with these sera (1:50) (after blocking aspecific protein-binding sites). The human tuberculous sera are selected for their reactivity (high or low) against the purified 32-kDa antigen of BCG tested in a dot blot assay as described document (31) of the bibliography hereafter. Reactive areas on the nitrocellulose sheets revealed by incubation with peroxidase conjugated goat anti-human immunoglobulin G antibody (Dakopatts, Copenhagen, Denmark) (1:200) for 4h, and after repeated washings, color reaction is developed by adding (α-chloronaphtol) (Bio-Rad peroxidase substrate Laboratories, Richmond, Calif.) in the presence of peroxidase and hydrogen peroxide.

It goes without saying that the free reactive functions which are present in some of the amino acids, which are part of the constitution of the polypeptides of the invention, particularly the free carboxyl groups which are carried by the groups Glu or by the C-terminal amino acid on the one hand and/or the free NH₂ groups carried by the N-terminal amino acid or by amino acid inside the peptidic chain, for instance Lys, on the other hand, can be modified in so far as this modification does not alter the above mentioned properties of the polypeptide.

The molecules which are thus modified are naturally part of the invention. The above mentioned carboxyl groups can be acylated or esterified.

Other modifications are also part of the invention. Particularly, the amine or ester functions or both of terminal amino acids can be themselves involved in the bond with other amino acids. For instance, the N-terminal amino acid can be linked to a sequence comprising from 1 to several amino acids corresponding to a part of the C-terminal region of another peptide.

Furthermore, any peptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids of the polypeptides according to the invention are part of the invention in so far as this modification does not alter the above mentioned properties of said polypeptides.

The polypeptides according to the invention can be glycosylated or not, particularly in some of their glycosylation sites of the type Asn-X-Ser or Asn-X-Thr, X representing any amino acid.

Advantageous recombinant polypeptides of the invention contain in their polypeptidic chain, one at least of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-49) to to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b.

Advantageous recombinant polypeptides of the invention contain in their polypeptidic chain, one at least of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-49) to to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b.

Advantageous recombinant polypeptides of the invention contain in their polypeptidic chain, one at least of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-43) to the extremity constituted by amino acid at position (-1) represented on fig. 5.

Advantageous recombinant polypeptides of the invention contain in their polypeptidic chain, one at least of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity

constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,

- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b.

Advantageous recombinant polypeptides of the invention contain in their polypeptidic chain, one at least of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-42) to the extremity

constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,

- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b.

Advantageous recombinant polypeptides of the invention contain in their polypeptidic chain, one at least of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (295) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-30) to the extremity constituted by amino acid at position (295) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-43) to the extremity constituted by amino acid at position (295) represented on fig. 5.

Other advantageous recombinant polypeptides of the invention consist in one of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b.

Other advantageous recombinant polypeptides of the invention consist in one of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,

- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b.

Other advantageous recombinant polypeptides of the invention consist in one of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (295) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-30) to the extremity constituted by amino acid at position (295) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-43) to the extremity

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constituted by amino acid at position (295) represented on fig. 5.

Other advantageous recombinant polypeptides of the invention consist in one of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b.

Other advantageous recombinant polypeptides of the invention consist in one of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,

- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b.

Other advantageous recombinant polypeptides of the invention consist in one of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-43) to the extremity constituted by amino acid at position (-1) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-30) to the extremity constituted by amino acid at position (-1) represented on fig. 5.

In eukaryotic cells, these polypeptides can be used as signal peptides, the role of which is to initiate the translocation of a protein from its site of synthesis, but which is excised during translocation.

Other advantageous peptides of the invention consist in one of the following amino acid sequence:

- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (275) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b.

Other advantageous peptides of the invention consist in one of the following amino acid sequence:

- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 4a and fig. 4b, or

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- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (275) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b.

Other advantageous peptides of the invention consist in one of the following amino acid sequence:

- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (77) to the extremity

constituted by amino acid at position (96) represented on fig. 5, or

- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (275) to the extremity constituted by amino acid at position (295) represented on fig. 5.

It is to be noted that the above mentioned polypeptides are derived from the expression products of a DNA derived from the nucleotide sequence coding for a protein of 32-kDa secreted by Mycobacterium tuberculosis as explained hereafter in the examples.

The invention also relates to the amino acid sequences constituted by the above mentioned polypeptides and a protein or an heterologous sequence with respect to said polypeptide, said protein or heterologous sequence comprising for instance from about 1 to about 1000 amino acids. These amino acid sequences will be called fusion proteins.

In an advantageous fusion protein of the invention, the heterologous protein is β -galactosidase.

Other advantageous fusion proteins of the invention are the ones containing an heterologous protein resulting from the expression of one of the following plasmids:

pEX1
pEX2
pEX3
pUEX1 pmTNF MPH
pUEX2
pUEX3

The invention also relates to any nucleotide sequence coding for a polypeptide of the invention.

The invention also relates to nucleic acids comprising nucleotide sequences which hybridize with the nucleotide sequences coding for any of the above mentioned polypeptides under the following hybridization conditions:

- hybridization and wash medium: 3 X SSC, 20% formamide (1 X SSC is 0,15 M NaCl, 0.015 M sodium citrate, pH 7.0),
- hybridization temperature (HT) and wash temperature (WT) for the nucleic acids of the invention defined by x-y: i.e. by the sequence extending from the extremity constituted by the nucleotide at position (x) to the extremity constituted by the nucleotide at position (y) represented on fig. 3a and fig. 3b.
 - 1 182 $HT = WT = 69 \cdot C$ 1 -194 $HT = WT = 69 \cdot C$ 1 -212 $HT = WT = 69 \cdot C$ $HT = WT = 69 \cdot C$ 1 -218 1 - 272 $HT = WT = 69 \cdot C$ 1 -359 $HT = WT = 71^{\circ}C$ 1 - 1241 $HT = WT = 73 ^{\circ}C$ 1 - 1358 $HT = WT = 73 \cdot C$ $HT = WT = 70^{\circ}C$ 183 -359 $HT = WT = 73 \cdot C$ 183 - 1241 183 - 1358 $HT = WT = 73 \cdot C$ $HT = WT = 70^{\circ}C$ 195 - 359 195 - 1241 $HT = WT = 73 \cdot C$ 195 - 1358 $HT = WT = 73 \cdot C$ $HT = WT = 70^{\circ}C$ 213 - 359

213	-	1241	HT	=	WT	=	73°C
213	_	1358	HT	=	WT	=	73°C
219	_	359	HT	=	WT	=	71°C
219	_	1241	HT	=	WT	==	73°C
219	-	1358	HT	=	WT	=	73°C
234	_	359	HT	=	WT	=	71°C
234	-	1241	HT	=	WT	=	74°C
234	_	1358	HT	=	WT	=	73°C
273	-	359	HT	=	WT	=	71°C
273	-	1241	HT	=	WT	=	74°C
273	_	1358	HT	=	WT	=	73°C
360	-	1241	HT	=	WT	=	73°C
360	-	1358	HT	=	WT	=	73°C
1242	-	1358	HT	=	WT	=	62°C

The above mentioned temperatures are to be considered as approximately ± 5°C.

The invention also relates to nucleic acids comprising nucleotide sequences which are complementary to the nucleotide sequences coding for any of the above mentioned polypeptides.

It is to be noted that in the above defined nucleic acids, as well as in the hereafter defined nucleic acids, the nucleotide sequences which are brought into play are such that T can be replaced by U.

A group of preferred nucleic acids of the invention comprises one at least of the following nucleotide sequences:

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (182) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1242) to the extremity constituted by nucleotide at position (1358), wherein N represents one of the five A, T, C, G or I nucleotides, represented in fig. 3a and fig. 3b,
- or above said nucleotide sequences wherein T is replaced by U,
- or nucleic acids which hybridize with said above mentioned nucleotide sequences or the complementary sequences thereof.

A group of preferred nucleic acids of the invention comprises one at least of the following nucleotide sequences:

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (182) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1242) to the extremity constituted by nucleotide at position (1358), wherein N represents one of the five A, T, C, G or I nucleotides, represented in fig. 4a and fig. 4b,
- or above said nucleotide sequences wherein ${\bf T}_{\cdot}$ is replaced by ${\bf U}_{\star}$

or nucleic acids which hybridize with said above mentioned nucleotide sequences or the complementary sequences thereof.

A group of preferred nucleic acids of the invention comprises one at least of the following nucleotide sequences:

- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (219) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (220) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1104) to the extremity constituted by nucleotide at position (1299), wherein N represents one of the five A, T, C, G or I nucleotides, represented in fig. 5,
- or above said nucleotide sequences wherein T is replaced by U,
- or nucleic acids which hybridize with said above mentioned nucleotide sequences or the complementary sequences thereof.

Other preferred nucleic acids of the invention comprise one at least of the following nucleotide sequences:

- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b.

Other preferred nucleic acids of the invention comprise one at least of the following nucleotide sequences:

- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b.

Another preferred group of nucleic acids of the invention comprises the following nucleotide sequences: - the one extending from the extremity constituted by nucleotide at position (360) to the constituted by nucleotide at position (1358)represented in fig. 3a and fig. 3b.

Another preferred group of nucleic acids of the invention comprises the following nucleotide sequences:

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- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b.

According to another advantageous embodiment, nucleic acids of the invention comprises one of the following sequences:

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (194) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (212) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (218) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (272) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,

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- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b.

According to another advantageous embodiment, nucleic acids of the invention comprises one of the following sequences:

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (194) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (212) represented in fig. 4a and fig. 4b.
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (218) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (272) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,

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- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b.

- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity

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constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b.

- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b.

These nucleotide sequence can be used as nucleotide signal sequences, coding for the corresponding signal peptide.

Preferred nucleic acids of the invention consist in one of the following nucleotide sequences:

- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b.

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (182) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (194) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (212) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (218) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (272) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1242) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b.

- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (360) to the extremity

constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b.

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (182) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (194) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (212) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (218) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (272) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,

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- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,

- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1242) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b.

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (129) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (219) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (90) to the extremity constituted by nucleotide at position (219) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (90) to the extremity

constituted by nucleotide at position (1299) represented in fig. 5,

- the one extending from the extremity constituted by nucleotide at position (90) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (220) to the extremity constituted by nucleotide at position (1299) represented in fig. 5.

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (129) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (219) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (90) to the extremity constituted by nucleotide at position (219) represented in fig. 5,

- the one extending from the extremity constituted by nucleotide at position (90) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (90) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (219) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (220) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (220) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1104) to the extremity constituted by nucleotide at position (1299) represented in fig. 5.

The invention also relates to any recombinant nucleic acids containing at least a nucleic acid of the invention inserted in an heterologous nucleic acid.

The invention relates more particularly to recombinant nucleic acid such as defined, in which the nucleotide sequence of the invention is preceded by a promoter (particularly an inducible promoter) under the control of which the transcription of said sequence is liable to be processed and possibly followed by a sequence coding for transcription termination signals.

The invention also relates to the recombinant nucleic acids in which the nucleic acid sequences coding for the polypeptide of the invention are recombined with peptide, signal possibly the control elements which are heterologous with respect to the ones to which they are normally associated within particularly, more and. bacteria gene the regulation elements adapted to control their expression in the cellular host which has been chosen for their production.

The invention also relates to recombinant vectors, particularly for cloning and/or expression, comprising a vector sequence, notably of the type plasmid, cosmid or phage, and a recombinant nucleic acid of the invention, in one of the non essential sites for its replication.

Appropriate vectors for expression of the recombinant antigen are the following one:

pEX1 pmTNF MPH
pEX2 pIGRI
pEX3
pUEX1
pUEX2
pUEX3

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The pEX1, pEX2 and pEX3 vectors are commercially available and can be obtained from Boehringer Mannheim.

The pUEX1, pUEX2 and pUEX3 vectors are also commercially available and can be obtained from Amersham.

According to an advantageous embodiment of the invention, the recombinant vector contains, in one of its non essential sites for its replication, necessary elements to promote the expression of polypeptides according to the invention in a cellular host and possibly a promoter recognized by the polymerase of the cellular host, particularly an inducible promoter and possibly a signal sequence and/or an anchor sequence.

According to another additional embodiment of the invention, the recombinant vector contains the elements enabling the expression by $\underline{\mathbf{E.}}$ coli of a nucleic acid according to the invention inserted in the vector, and particularly the elements enabling the expression of the gene or part thereof of β -galactosidase.

The invention also relates to a cellular host which is transformed by a recombinant vector according to the invention, and comprising the regulation elements enabling the expression of the nucleotide sequence coding for the polypeptide according to the invention in this host.

The invention also relates to a cellular host chosen from among bacteria such as <u>E. coli</u>, transformed by a vector as above defined, and defined hereafter in the examples, or chosen from among eukaryotic organism, such as CHO cells, insect cells, Sf9 cells [Spodoptera frugiperda] infected by the virus Ac NPV (Autographa californica nuclear polyhydrosis virus) containing suitable vectors such as pAc 373 pYM1 or pVC3, BmN [Bombyx mori] infected by the virus BmNPV containing suitable vectors such as pBE520 or p89B310.

The invention relates to an expression product of a nucleic acid expressed by a transformed cellular host according to the invention.

The invention also relates to nucleotidic probes, hybridizing with anyone of the nucleic acids or with their complementary sequences,

and particularly the probes chosen among the following nucleotidic sequences gathered in Table 1, and represented in fig. 9.

TABLE 1

Probes A(i), A(ii), A(iii), A(iv) and A(v)

- A(i) CAGCTTGTTGACAGGGTTCGTGGC
- A(ii) GGTTCGTGGCGCCGTCACG
- A(iii) CGTCGCGCGCCTAGTGTCGG
- A(iv) CGGCGCCGTCGGTGGCACGGCGA
- A(v) CGTCGGCGCGCCCTAGTGTCGG

Probe B

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TCGCCCGCCCTGTACCTG

Probe C

GCGCTGACGCTGGCGATCTATC

Probe D

CCGCTGTTGAACGTCGGGAAG

Probe E

AAGCCGTCGGATCTGGGTGGCAAC

Probes F(i), F(ii), F(iii) and F(iv)

- F(i) ACGGCACTGGGTGCCACGCCCAAC
- F(ii) ACGCCCAACACCGGGCCCGCCA
- F(iii) ACGGGCACTGGGTGCCACGCCCAAC
- F(iv) ACGCCCCAACACCGGGCCCGCGCCCCA
- or their complementary nucleotidic sequences.

The hybridization conditions can be the following ones:

- hybridization and wash medium: 3 X SSC, 20% formamide (1 X SSC is 0,15 M NaCl, 0.015 M sodium citrate,

pH 7.0),

- hybridization temperature (HT) and wash temperature (WT):

(WT) °C:	HT and WT (°C)
A(i)	50
A(ii)	50
A(iii)	52
A(iv)	60
A(v)	52
В	48
С	50
D	45
E	52
F(i)	55
F(ii)	59
F(iii)	55
F(iv)	59

These probes might enable to differentiate <u>M.</u>

<u>tuberculosis</u> from other bacterial strains and in particular from the following mycobacteria species:

- Mycobacterium marinum, Mycobacterium scrofulaceum, Mycobacterium gordonae, Mycobacterium szulgai, Mycobacterium intracellulare, Mycobacterium xenopi, Mycobacterium gastri, Mycobacterium nonchromogenicum, Mycobacterium terrae and Mycobacterium triviale, and more particularly from \underline{M} . \underline{bovis} , Mycobacterium kansasii, Mycobacterium avium, Mycobacterium phlei and Mycobacterium fortuitum.

The invention also relates to DNA or RNA primers which can be used for the synthesis of nucleotidic sequences according to the invention by PCR (polymerase chain reaction technique), such as described in US Patents n° 4,683,202 and n° 4,683,195 and European Patent n° 200362.

The invention also relates to any DNA or RNA primer constituted by about 15 to about 25 nucleotides

of a nucleotide sequence coding for a polypeptide according to the invention.

The invention also relates to any DNA or RNA primer constituted by about 15 to about 25 nucleotides liable to hybridize with a nucleotide sequence coding for a polypeptide according to the invention.

The invention also relates to any DNA or RNA primer constituted by about 15 to about 25 nucleotides complementary to a nucleotide sequence coding for a polypeptide according to the invention.

The sequences which can be used as primers are given in Table 2 hereafter (sequences P1 to P6 or their complement) and illustrated in fig. 9:

TABLE 2

P1 GAGTACCTGCAGGTGCCGTCGCCGTCGATGGGCCG

P2 ATCAACACCCGGCGTTCGAGTGGTAC

P2 compl. GTACCACTCGAACGCCGGGGTGTTGAT

P3 TGCCAGACTTACAAGTGGGA

P3 compl. TCCCACTTGTAAGTCTGGCA

P4 TCCTGACCAGCGAGCTGCCG

P4 compl. CGGCAGCTCGCTGGTCAGGA

P5 CCTGATCGGCCTGGCGATGGGTGACGC

P5 compl. GCGTCACCCATCGCCAGGCCGATCAGG

P6 compl. GCGCCCAGTACTCCCAGCTGTGCGT

compl. = complement

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The sequences can be combined in twelve different primer-sets (given in Table 3) which allow enzymatical amplification by the polymerase chain reaction (PCR) technique of any of the nucleotide sequences of the invention, and more particularly the one extending from the extremity constituted by nucleotide at position 1 to the extremity constituted by nucleotide at position 1358, as well as the nucleotide sequence of antigen α of BCG (17).

The detection of the PCR amplified product can be achieved by a hybridization reaction with an oligonucleotide sequence of at least 10 nucleotides which is located between PCR primers which have been used to amplify the DNA.

The PCR products of the nucleotide sequences of the invention can be distinguished from the α -antigen gene of BCG or part thereof by hybridization techniques (dot-spot, Southern blotting, etc.) with the probes indicated in Table 3. The sequences of these probes can be found in Table 1 hereabove.

TABLE 3

Prime	er s	set					Detection with probe
1.	P1	and	the	complement	of	P2	В
2.	P1	and	the	complement	of	P3	В
3.	P1	and	the	complement	of	P4	В
4.	P1	and	the	complement	of	P 5	B or C
5.	Pl	and	the	complement	of	P6	B, C, D or E
6.	P2	and	the	complement	of	P 5	C
7.	P2	and	the	complement	of	P6	C, D or E
8.	P3	and	the	complement	of	P5	c .
9.	P 3	and	the	complement	of	P6	C, D or E
10.	P4	and	the	complement	of	P5	c
11.	P4	and	the	complement	of	Pé	C, D or E
12.	P5	and	the	complement	of	Pé	D or E

It is to be noted that enzymatic amplification can also be achieved with all oligonucleotides with sequences of about 15 consecutive bases of the primers given in Table 2. Primers with elongation at the 5'-end or with a small degree of mismatch may not considerably affect the outcome of the enzymatic amplification if the mismatches do not interfere with the base-pairing at the 3'-end of the primers.

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Specific enzymatic amplification of the nucleotide sequences of the invention and not of the BCG gene can be achieved when the probes (given in Table 1) or their complements are used as amplification primers.

When the above mentioned probes of Table 1 are used as primers, the primer sets are constituted by any of the nucleotide sequences (A, B, C, D, E, F) of Table 1 in association with the complement of any other nucleotide sequence, chosen from A, B, C, D, E or F, it being understood that sequence A means any of the sequences A(i), A(ii), A(iii), A(iv), A(v) and sequence F, any of the sequences F(i), F(ii), F(iii) and F(iv).

Advantageous primer sets for enzymatic amplification of the nucleotide sequence of the invention can be one of the following primer sets given in Table 3bis hereafter:

TABLE 3BIS

	A(i)					
or	A(ii)					
or	A(iii)	and	the	complement	of	В
or	A(iv)					
or	A(V)					
	A(i)					
or	A(ii)					
or	A(iii)	and	the	complement	of	С
or	A(iv)					
or	A(V)					
	В	and	the	complement	of	С
	A(i)					
or	A(ii)					
or	A(iii)	and	the	complement	of	F
or	A(iv)					
or	A(v)					

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A(i)
or A(ii)
                 and the complement of D
or A(iii)
or A(iv)
or A(v)
   A(i)
or A(ii)
                  and the complement of E
or A(iii)
or A(iv)
or A(V)
                   and the complement of D
   В
                   and the complement of E
   В
                   and the complement of F
   В
                   and the complement of D
   C
                   and the complement of E
   C
                   and the complement of F
   C
                   and the complement of E
   D
                   and the complement of F
   D
                   and the complement of F
   E
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A(i), A(ii), A(iii), A(iv), A(v), B, C, D, E and F having the nucleotide sequence indicated in Table 1.

In the case of amplification of nucleotide a sequence of the invention with any of the above mentioned primer sets defined in Table 3bis hereabove, the detection of the amplified nucleotide sequence can a hybridization reaction with an be achieved by oligonucleotide sequence of at least 10 nucleotides, said sequence being located between the PCR primers which have been used to amplify the nucleotide sequence. An oligonucleotide sequence located between said two primers can be determined from figure 9 where the primers A, B, C, D, E and F are represented by the boxed sequences respectively named probe region A, probe region B, probe region C, probe region D, probe region E and probe region F.

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The invention also relates to a kit for enzymatic amplification of a nucleotide sequence by PCR technique and detection of the amplified nucleotide sequence containing

- one of the PCR primer sets defined in Table 3 and one of the detection probes of the invention, advantageously the probes defined in Table 1,

or one of the PCR primer sets defined in Table 3bis, and a detection sequence consisting for instance in an oligonucleotide sequence of at least 10 nucleotides, said sequence being located (fig. 9) between the two PCR primers constituting the primer set which has been used for amplifying said nucleotide sequence.

The invention also relates to a process for preparing a polypeptide according to the invention comprising the following steps:

- the culture in an appropriate medium of a cellular host which has previously been transformed by an appropriate vector containing a nucleic acid according to the invention,
- the recovery of the polypeptide produced by the above said transformed cellular host from the above said culture medium, and
- the purification of the polypeptide produced, eventually by means of immobilized metal ion affinity chromatography (IMAC).

The polypeptides of the invention can be prepared according to the classical techniques in the field of peptide synthesis.

The synthesis can be carried out in homogeneous solution or in solid phase.

For instance, the synthesis technique in homogeneous solution which can be used is the one described by Houbenweyl in the book titled "Methode der organischen chemie" (Method of organic chemistry)

edited by E. Wunsh, vol. 15-I et II. THIEME, Stuttgart 1974.

The polypeptides of the invention can also be prepared according to the method described by R.D. MERRIFIELD in the article titled "Solid phase peptide synthesis" (J.P. Ham.Socks., 45, 2149-2154).

The invention also relates to a process for preparing the nucleic acids according to the invention.

A suitable method for chemically preparing the single-stranded nucleic acids (containing at most 100 nucleotides of the invention) comprises the following steps:

- DNA synthesis using the automatic β -cyanoethyl phosphoramidite method described in Bioorganic Chemistry 4; 274-325, 1986.

In the case of single-stranded DNA, the material which is obtained at the end of the DNA synthesis can be used as such.

A suitable method for chemically preparing the double-stranded nucleic acids (containing at most 100 bp of the invention) comprises the following steps:

- DNA synthesis of one sense oligonucleotide using the automatic β -cyanoethyl phosphoramidite method described in Bioorganic Chemistry 4; 274-325, 1986, and DNA synthesis of one anti-sense oligonucleotide using said above-mentioned automatic β -cyanoethyl phosphoramidite method,
- combining the sense and anti-sense oligonucleotides by hybridization in order to form a DNA duplex,
- cloning the DNA duplex obtained into a suitable plasmid vector and recovery of the DNA according to classical methods, such as restriction enzyme digestion and agarose gel electrophoresis.

A method for the chemical preparation of nucleic acids of length greater than 100 nucleotides - or bp,

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in the case of double-stranded nucleic acids - comprises the following steps:

- assembling of chemically synthesized oligonucleotides, provided at their ends with different restriction sites, the sequences of which are compatible with the succession of amino acids in the natural peptide, according to the principle described in Proc. Nat. Acad. Sci. USA 80; 7461-7465, 1983,
- cloning the DNA thereby obtained into a suitable plasmid vector and recovery of the desired nucleic acid according to classical methods, such as restriction enzyme digestion and agarose gel electrophoresis.

The invention also relates to antibodies themselves formed against the polypeptides according to the invention.

It goes without saying that this production is not limited to polyclonal antibodies.

It also relates to any monoclonal antibody produced by any hybridoma liable to be formed according to classical methods from splenic cells of an animal, particularly of a mouse or rat, immunized against the purified polypeptide of the invention on the one hand, and of cells of a myeloma cell line on the other hand, and to be selected by its ability to produce the monoclonal antibodies recognizing the polypeptide which has been initially used for the immunization of the animals.

The invention also relates to any antibody of the invention labeled by an appropriate label of the enzymatic, fluorescent or radioactive type.

The peptides which are advantageously used to produce antibodies, particularly monoclonal antibodies, are the following ones gathered in Table 4:

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TABLE 4a (see fig. 4a and 4b)

Amino acid		Amino acid
position		position
$(NH_2-terminal)$		(COOH-terminal)
12	QVPSPSMGRDIKVQFQSGGA	31
36	LYLLDGLRAQDDFSGWDINT	55
77	SFYSDWYQPACRKAGCQTYK	96
101	LTSELPGWLQANRHVKPTGS	. 120
175	KASDMWGPKEDPAWQRNDPL	194
211	CGNGKPSDLGGNNLPAKFLE	230
275	KPDLQRHWVPRPTPGPPQGA	294

TABLE 4b (see fig. 5)

Amino acid	1	Amino acid
position		position
(NH ₂ -termi	nal)	(COOH-terminal)
77	SFYSDWYQPACGKAGCQTYK	96
276	PDLQRALGATPNTGPAPQGA	295

The amino acid sequences are given in the 1-letter code.

Variations of the peptides listed in Table 4 are also possible depending on their intended use. For example, if the peptides are to be used to raise antisera, the peptides may be synthesized with an extra cysteine residue added. This extra cysteine residue is preferably added to the amino terminus and facilitates the coupling of the peptide to a carrier protein which is necessary to render the small peptide immunogenic. If the peptide is to be labeled for use in radioimmune assays, it may be advantageous to synthesize the protein with a tyrosine attached to either the amino or carboxyl terminus to facilitate iodination. These

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peptides possess therefore the primary sequence of the peptides listed in Table 4 but with additional amino acids which do not appear in the primary sequence of the protein and whose sole function is to confer the desired chemical properties to the peptides.

The invention also relates to a process for detecting in vitro antibodies related to tuberculosis in a human biological sample liable to contain them, this process comprising

- contacting the biological sample with a polypeptide or a peptide according to the invention under conditions enabling an <u>in vitro</u> immunological reaction between said polypeptide and the antibodies which are possibly present in the biological sample and
- the <u>in vitro</u> detection of the antigen/antibody complex which may be formed.

Preferably, the biological medium is constituted by a human serum.

The detection can be carried out according to any classical process.

By way of example a preferred method brings into play an immunoenzymatic process according to ELISA technique or immunofluorescent or radioimmunological (RIA) or the equivalent ones.

Thus the invention also relates to any polypeptide according to the invention labeled by an appropriate label of the enzymatic, fluorescent, radioactive... type.

Such a method for detecting <u>in vitro</u> antibodies related to tuberculosis comprises for instance the following steps:

- deposit of determined amounts of a polypeptidic composition according to the invention in the wells of a titration microplate,
- introduction into said wells of increasing dilutions of the serum to be diagnosed,

- incubation of the microplate,
- repeated rinsing of the microplate,
- introduction into the wells of the microplate of labeled antibodies against the blood immunoglobulins,
- the labeling of these antibodies being carried out by means of an enzyme which is selected from among the ones which are able to hydrolyze a substrate by modifying the absorption of the radiation of this latter at least at a given wave length,
- detection by comparing with a control standard of the amount of hydrolyzed substrate.

The invention also relates to a process for detecting and identifying in vitro antigens of M. tuberculosis in a human biological sample liable to contain them, this process comprising:

- contacting the biological sample with an appropriate antibody of the invention under conditions enabling an <u>in vitro</u> immunological reaction between said antibody and the antigens of <u>M. tuberculosis</u> which are possibly present in the biological sample and the <u>in vitro</u> detection of the antigen/antibody complex which may be formed.

Preferably, the biological medium is constituted by sputum, pleural effusion liquid, broncho-alveolar washing liquid, urine, biopsy or autopsy material.

Appropriate antibodies are advantageously monoclonal antibodies directed against the peptides which have been mentioned in Table 4.

The invention also relates to an additional method for the <u>in vitro</u> diagnostic of tuberculosis in a patient liable to be infected by Mycobacterium tuberculosis comprising the following steps:

- the possible previous amplification of the amount of the nucleotide sequences according to the invention, liable to be contained in a biological sample taken \$

from said patient by means of a DNA primer set as above defined,

- contacting the above mentioned biological sample with a nucleotide probe of the invention, under conditions enabling the production of an hybridization complex formed between said probe and said nucleotide sequence, - detecting the above said hybridization complex which has possibly been formed.

To carry out the <u>in vitro</u> diagnostic method for tuberculosis in a patient liable to be infected by Mycobacterium tuberculosis as above defined, the following necessary or kit can be used, said necessary or kit comprising:

- a determined amount of a nucleotide probe of the invention,
- advantageously the appropriate medium for creating an hybridization reaction between the sequence to be detected and the above mentioned probe,
- advantageously, reagents enabling the detection of the hybridization complex which has been formed between the nucleotide sequence and the probe during the hybridization reaction.

The invention also relates to an additional method for the <u>in vitro</u> diagnostic of tuberculosis in a patient liable to be infected by Mycobacterium tuberculosis comprising:

- contacting a biological sample taken from a patient with a polypeptide or a peptide of the invention, under conditions enabling an <u>in vitro</u> immunological reaction between said polypeptide or peptide and the antibodies which are possibly present in the biological sample and - the <u>in vitro</u> detection of the antigen/antibody complex which has possibly been formed.

To carry out the $\underline{\text{in}}$ $\underline{\text{vitro}}$ diagnostic method for tuberculosis in a patient liable to be infected by

Mycobacterium tuberculosis, the following necessary or kit can be used, said necessary or kit comprising:

- a polypeptide or a peptide according to the invention.
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complex which has been produced by the immunological reaction, said reagents possibly having a label, or being liable to be recognized by a labeled reagent, more particularly in the case where the above mentioned polypeptide or peptide is not labeled.

The invention also relates to an additional method for the <u>in vitro</u> diagnostic of tuberculosis in a patient liable to be infected by <u>M. tuberculosis</u>, comprising the following steps:

- contacting the biological sample with an appropriate antibody of the invention under conditions enabling an <u>in vitro</u> immunological reaction between said antibody and the antigens of <u>M. tuberculosis</u> which are possibly present in the biological sample and - the <u>in vitro</u> detection of the antigen/antibody complex which may be formed.

Appropriate antibodies are advantageously monoclonal antibodies directed against the peptides which have been mentioned in Table 4.

To carry out the <u>in vitro</u> diagnostic method for tuberculosis in a patient liable to be infected by Mycobacterium tuberculosis, the following necessary or kit can be used, said necessary or kit comprising:

- an antibody of the invention,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complexes which have been produced by the immunological reaction, said reagent possibly having a label or being

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liable to be recognized by a label reagent, more particularly in the case where the above mentioned antibody is not labeled.

An advantageous kit for the diagnostic <u>in vitro</u> of tuberculosis comprises:

- at least a suitable solid phase system, e.g. a microtiter-plate for deposition thereon of the biological sample to be diagnosed in vitro,

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- a preparation containing one of the monoclonal antibodies of the invention,
- a specific detection system for said monoclonal antibody,
- appropriate buffer solutions for carrying out the immunological reaction between a test sample and said monoclonal antibody on the one hand, and the bonded monoclonal antibodies and the detection system on the other hand.

The invention also relates to a kit, as described above, also containing a preparation of one of the polypeptides or peptides of the invention, said antigen of the invention being either a standard (for quantitative determination of the antigen of M. tuberculosis which is sought) or a competitor, with respect to the antigen which is sought, for the kit to be used in a competition dosage process.

The invention also relates to an immunogenic composition comprising a polypeptide or a peptide according to the invention, in association with a pharmaceutically acceptable vehicle.

The invention also relates to a vaccine composition comprising among other immunogenic principles anyone of the polypeptides or peptides of the invention or the expression product of the invention, possibly coupled to a natural protein or to a synthetic polypeptide having a sufficient molecular weight so that the conjugate is able to induce in vivo

the production of antibodies neutralizing Mycobacterium tuberculosis, or induce <u>in vivo</u> a cellular immune response by activating M. tuberculosis antigenresponsive T cells.

The peptides of the invention which are advantageously used as immunogenic principle have one of the following sequences:

TABLE 4a (see fig. 4a and 4b)

Amino acid		Amino acid
position		position
(NH ₂ -terminal)		(COOH-terminal)
12	QVPSPSMGRDIKVQFQSGGA	31
36	LYLLDGLRAQDDFSGWDINT	55
77	SFYSDWYQPACRKAGCQTYK	96
101	LTSELPGWLQANRHVKPTGS	120
175	KASDMWGPKEDPAWQRNDPL	194
211	CGNGKPSDLGGNNLPAKFLE	230
275	KPDLQRHWVPRPTPGPPQGA	294

TABLE 4b (see fig. 5)

Amino acid		Amino acid
position		position
(NH ₂ -terminal)	(COOH-terminal)
77	SFYSDWYOPACGKAGCOTYK	96
276	PDLQRALGATPNTGPAPQGA	299

The amino acid sequences are given in the 1-letter code.

Other characteristics and advantages of the invention will appear in the following examples and the figures illustrating the invention.

Figures 1(A) and 1(B) correspond to the identification of six purified $\lambda gt11$ M. tuberculosis recombinant clones. Figure 1(A) corresponds to the

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EcoRI restriction analysis of clone 15, clone 16, clone 17, clone 19, clone 24 and EcoRI-HindIII digested lambda DNA-molecular weight marker lane (in kilobase pairs) (M) (Boehringer).

Figure 1(B) corresponds to the immunoblotting analysis of crude lysates of E. coli lysogenized with clone 15, clone 16, clone 17, clone 19, clone 23 and clone 24.

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Arrow (<—) indicates fusion protein produced by recombinant λ gtll-M-tuberculosis clones. Expression and immunoblotting were as described above. Molecular weight (indicated in kDa) were estimated by comparison with molecular weight marker (High molecular weight-SDS calibration kit, Pharmacia).

Figure 2 corresponds to the restriction map of the DNA inserts in the λgt11 M. tuberculosis recombinant clones 17 and 24 identified with polyclonal anti-32-kDa (BCG) antiserum as above defined and of clones By1, By2 and By5 selected by hybridization with a 120 bp EcoRI-Kpn I restriction fragment of clone 17.

DNA was isolated from Agt11 phage stocks by using the Lambda Sorb Phage Immunoadsorbent, as described by the manufacturer (Promega). Restriction sites were located as described above. Some restriction sites (*) were deduced from a computer analysis of the nucleotide sequence.

The short vertical bars (|---|) represent linker derived EcoRI sites surrounding the DNA inserts of recombinant clones. The lower part represents a magnification of the DNA region containing the antigen of molecular weight of 32-kDa, that has been sequenced. Arrows indicate strategies and direction of dideoxy-sequencing. (--->) fragment subcloned in Bluescribe M13; (<-->) fragment subcloned in mp10 and mp11 M13 vectors; (--->) sequence determined with the use of a synthetic oligonucleotide.

Figures 3a and 3b correspond to the nucleotide and amino acid sequences of the general formula of the antigens of the invention.

Figures 4a and 4b correspond to the nucleotide and amino acid sequences of one of the antigens of the invention.

Two groups of sequences resembling the E. coli consensus promoter sequences are boxed and the homology to the consensus is indicated by italic bold letters. Roman bold letters represent a putative Shine-Dalgarno motif.

Figure 5 corresponds to the nucleotide and amino acid sequences of the antigen of 32-kDa of the invention.

The NH_2 -terminal amino acid sequence of the mature protein which is underlined with a double line happens to be very homologous - 29/32 amino acids - with the one of MPB 59 antigen (34). Vertical arrows (ψ) indicate the presumed NH_2 end of clone 17 and clone 24.

Figure 6 is the hydropathy pattern of the antigen of the invention of a molecular weight of 32-kDa and of the antigen α of BCG (17).

Figure 7 represents the homology between the amino acid sequences of the antigen of 32-kDa of the invention and of antigen α of BCG (revised version).

Identical amino acids; (:) evolutionarily conserved replacement of an amino acid (.), and absence

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of homology () are indicated. Underlined sequence (=) represents the signal peptide, the option taken here arbitrarily representing the 43-amino acid signal peptide corresponding to ATG₉₁. Dashes in the sequences indicate breaks necessary for obtaining the optimal alignment.

Figure 8 illustrates the fact that the protein of 32-kDa of the invention is selectively recognized by human tuberculous sera.

Figure 8 represents the immunoblotting with human tuberculous sera, and anti- β -galactosidase monoclonal antibody. Lanes 1 to 6: E. coli lysate expressing fusion protein (140 kDa); lanes 7 to 12:unfused β -galactosidase (114 kDa). The DNA insert of clone 17 (2.7 kb) was subcloned into pUEX2 and expression of fusion protein was induced as described by Bresson and Stanley (4). Lanes 1 and 7 were probed with the anti- β -galactosidase monoclonal antibody: lanes 4, 5, 6 and 10, 11, 12 with 3 different human tuberculous sera highly responding towards purified protein of the invention of 32-kDa; lanes 2 and 3 and 8 and 9 were probed with 2 different low responding sera.

Figure 9 represents the nucleic acid sequence alignment of the 32-kDa protein gene of K. tuberculosis of the invention (upper line), corresponding to the sequence in fig. 5, of the gene of fig. 4a and 4b of the invention (middle line), and of the gene for antigen α of BCG (lower line).

Dashes in the sequence indicate breaks necessary for obtaining optimal alignment of the nucleic acid sequence.

The primer regions for enzymatical amplification are boxed (P1 to P6).

The specific probe regions are boxed and respectively defined by probe region A, probe region B,

probe region C, probe region D, probe region E and probe region F.

It is to be noted that the numbering of nucleotides is different from the numbering of figures 3a and figure 3b, and of figure 5, because nucleotide at position 1 (on figure 9) corresponds to nucleotide 234 on Figure 3a, and corresponds to nucleotide 91 on figure 5.

Figure 10a corresponds to the restriction and genetic map of the pIGRI plasmid used in Example IV for the expression of the P_{32} antigen of the invention in E. coli.

On this figure, underlined restriction sites are unique.

Figure 10b corresponds to the pIGRI nucleic acid sequence.

On this figure, the origin of nucleotide stretches used to construct plasmid pIGRI are specified hereafter.

Position

3422-206: lambda PL containing EcoRI blunt-MboII blunt fragment of $pPL(\lambda)$ (Pharmacia)

207-384 : synthetic DNA sequence

228-230: initiation codon ATG of first cistron

234-305: DNA encoding amino acids 2 to 25 of

mature mouse TNF

306-308: stop codon (TAA) first cistron

311-312: initiation codon (ATG) second

cistron

385-890: $rrnBT_1T_2$ containing HindIII-SspI

fragment from pKK223 (Pharmacia)

891-3421: DraI-EcoRI blunt fragment of pAT₁₅₃
(Bioexcellence) containing the

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tetracycline resistance gene and the origin of replication.

Table 5 hereafter corresponds to the complete restriction site analysis of pIGRI.

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Name of the plasmid : pIGRI

Total number of bases is: 3423. Analysis done on the complete sequence.

List of cuts by enzyme.

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3403	2529		
3388	2280		3202
2732	2156	2617	2813
2329	1977	2529	1869 2630
3003	1935	2280	1866 1997 3048
2982 1481	713	1977	1753 1973 1087 3035
2868 1345	623	1935	1735 1226 1001 2313
2765 2211 1088	494	494	1316 1223 2685 2487 903 999
370 735 1645 222 386 1236 1331	329 3244 1990	329 1973 3040 2214	389 1017 1822 2253 15 902
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	Nar I	Nco I	Nhe I	Nla III		Nla IV		Nra I				Ple I			Pss I						Sdu I		Sfa NI	Z	Sph I						Taq IIB*	Tth1111I	Tth111111*	Xba I	I		Xmn I

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Aat II Bgl II Eco RI Nsi I Sau I Spl I

List of

Sorted list of enzymes by n° of cuts.

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	, Bbv II , Bst XI , Nde I , Sac I , Sna BI , Tth 11
TthlllII' NSP BII Fok I Fok I Fok I Bok II Bsp HI Pss I Mst I Hgi JII Ple I Kou QI Acc I Bgl I Ple I Cvi QI Acc I Bgl I Flu III Rsa I Sal I Bbv II Bsp HI Sty I Bsp HI Sty I Bco NI	Avr II Bst EII Mme I Rsr II Sma I Taq IIA* f selected
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	selected Apa I Bep M Bep M Bep I Pou I Sci I Stu I Xma I
Cvi JI Fnu 461 Hha I Hha II Hha III Hha IV Nha III Rha III Scr FI Sco II Hho I	non cutting , Afl II' , Bsp MI* , Pma CI , Sca I , Ssp I , Xho I

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Figure 11a corresponds to the restriction and genetic map of the pmTNF MPH plasmid used in Example V for the expression of the P_{32} antigen of the invention in <u>E. coli</u>.

Figure 11b corresponds to the pmTNF-MPH nucleic acid sequence.

On this figure, the origin of nucleotide stretches used to construct plasmid pmTNF-MPH is specified hereafter.

Position

*

lambda PL containing EcoRI blunt-MboII 1-208: blunt fragment of $pPL(\lambda)$ (Pharmacia) synthetic DNA fragment 209-436: (ATG) of mTNF initiation codon 230-232 : fusion protein sequence encoding AA 2 to 25 of 236-307: mature mouse TNF multiple cloning site containing 308-384 : His, encoding sequence at position 315-332 HindIII fragment containing E. coli 385-436 : trp terminator

437-943: rrnBT₁T₂ containing HindIII-SspI fragment from pKK223 (Pharmacia)

944-3474: DraI-EcoRI blunt fragment of pAT $_{153}$ (Bioexcellence) containing the tetracycline resistance gene and the origin of replication.

Table 6 hereafter corresponds to the complete restriction site analysis of pmTNF-MPH.

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**********	* RESTRICTION-SITE ANALYSIS	****************

Done on DNA sequence PMTNFMPH.

Total number of bases is: 3474.
Analysis done on the complete sequence.

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List	

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P. A. J. M.	••	••	••	••	••	••	••	**	••	••	••		••	••	••	••	••	••	••	

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	3101					1552	1321	3067	i i		361	1141	1676	2215	2641	3175	1424	1040	3122									1550
	1140	3088				3	76	05	17		350	844	1658	2210	2597	3140	1015	962	3095									1537
	1054	2366				908	736	2699	3069		343	828	1632	2189	2552	3071	717	950	3004				3238			>		804
	95	05	97			3	4	53	93		272	167	1534	14	53	2999	661	336	2645	2945	6	1	2976		2000	9696	4	989
2738 2540	342	955	978			528	339	2185	2529		265	678	1398	0	S	2985 3306	571	238	1	3	2 0		2695		2022	2845	6	526
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Fnu DII	••	2447 542	2532 1074	2697 1655	2748 1837	2855 1934	2889 2056	2892 2082	3170 2227	3173 2237	3244 2366	243
Fok I Fok I* Gsu I	** ** **	2493 468 816 2088	2498 852 2423	2525 3370 2468	3322	2769	3125					
Gsu I* Hae I Hae II Hae III	** ** ** **	2642 361 594 343	828 1458 361	844 1828 678	1224 2267 767	1676 2697 828	1687 2924 844	2026 2978 1224	2423 3038 1658	2480 3059 1676	2552 3240 1687	202
		2210	2423	2480	2531	2552	2641	2875	2939	2947	3071	317
Hga I Hga I* Hgi AI Hgi CI	** ** ** **	3298 160 1008 141 210	183 1586 1388 2179	796 2482 2007 2263	2088 2514 2298 2702	2238 3068 2885 2920	2829 3196 3034	3055	3349	3392		
	• ••	542	593	1074	1183	1357	1457	1524	1794	1827	2017	205
Bin P1I	••	23 54 11	3371 591 2264	07	1181	1355 2694	1455 2769	1522 2921	1792 2975	1825 3035	2015	205
Hind II Hind III Hinf I	** ** **	3237 109 384 368	3369 372 437 1328	2819 3439 1724	1799	1944	2165	2463	2617	2837		

	m	6		~		6		=	9	1	6	
	198	330		232		25		201	346	22	33	
	1493	3083	3355	2008		2295		1881	3121	2181	3351	
	1346	3068	3073	3343 1132		2234	3436	1702	2983	2146	3141	
	1320	3059	3028	2569 3255 1057		2196	3414	982	2910	2032	3095	
	1130	2936	2174	2513 2587 1046	2743	2169	994	912	2764	1989	3057	
't)	169	2776	1953	1924 2320 1038	3120 2472	2683 1803	817	620	2725	1670	3036	
6 (Con't)	735	2700	196	3240 1900 1341 960	3093 1880	2499 1520	486	565	2539	1631	2946	
Table	375	2540	716	1981 997 1278 948	3002 1832	3252 2001 764	3347 388 3069 3056	382	2422	549	2922	2331
	355	2450	183 311	1205 751 1162 334	2643 970	3165 1595 350	3287 223 3210 2701 3035	349	2294	343	2704	1360 2910
	339	2212	140 305	952 330 257 236	2371 475	1489 1271 291	3083 188 2114 2541 2921	232	2279	336	2583	1115 1702
	ស	2186	3309 96 8	365 276 171 9	2340 209	1305 372 210	2864 181 2016 2187 2264	3239 168	2222	212	2265	2498 412 382
	••		•• •• ••		•• •		•• •• •• •	•• ••		••		
	II		* H H H			* * HHH	нннн	HHH		IV		I BII HI
	Нра		Hph Hph Hph	Mae Mae Mae Mae	Mbo	Mme Mn1 Mn1	Mse Mst Nae Nar	Nhe Nhe Nla		Nla		Nru Nsp Nsp

									153			335					153													
									1321			3339					1319													
									806	3340	3196	2940					804	3338			3446									
									169	3300	3001	2934					167	3298			3131									
									736	2936	2987	2301		3255			734	2934			2818						3093			
									638	2411	2885	2099	3344	3066			989	2409			2343						1057			
on't)									528	2212	2298	2021	3231	3054			526	2210			2202						1046			
Table 6 (Con't)									340	2028	2007	1538	2820	2433			338	2026			1600						960			
Tab	2154					2948			339	1986	1388	345	2445	2038			337	1984			999				1114		948			
	2105	1807	2831		2030	2033	3307	2817	215	1673	345	338	818	1601		2910	213	1671		2099	371			1107	1075		334			
	295	376	1322	331	1988	1991	212	370	9	1552	141	2	650	420	340	382	4	1550	361	345	254	1802	2804	40	989	364	6	338	2529	467
	••	••	••	••	••	••	••	••	••		••	••	••	••	••	••	••		••	••	••	••	••	••	••	••	••	••	••	••
	MI		* H								н	н	Z		Н	н			H	H	H	\mathbf{H}	IIB*	\vdash	11111*	H		H	III	н
	Pfl	Ple	Ple	Рта	Ppu	Pss	Rsa	Sal	Scr		Sdu	Sec	Sfa	Sfa	Sma	Sph	SBO		Stu	sty	Tag	Tag	Tag	Tth	Tthl	Xba	Xho	Хта	Хша	Xmn

Total number of cuts is

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Table 6 (con't)

List of non cutting selected enzymes.	su II , Avr II , Bbv II* , Bcl I , Bgl II , Bsp MI* st EII , Bst XI , Eco 311* , Esp I , Hpa I , Mlu I de I , Not I , Nsi I , Pst I , Pvu I , Pvu II ac I , Sac II , Sau I , Sca I , Sci I , Sfi I pe I , Spl I , Ssp I , Tag IIA , Tag IIA* , Tth 1111	of selected enzymes which do not cut: 38
List of non cutting sel	н	Total number of selecte

Figure 12a corresponds to the restriction and genetic map of the plasmid pIG2 used to make the intermediary construct pIG2 Mt32 as described in Example IV for the subcloning of the P_{32} antigen in plasmid pIGRI.

Figure 12b corresponds to the pIG2 nucleic acid sequence.

On this figure, the origin of nucleotide stretches used to construct plasmid pIG2 is specified hereafter.

Position

lambda PL containing EcoRI-MboII blunt 3300-206 : fragment of $pPL(\lambda)$ (Pharmacia) 207-266: synthetic sequence containing multiple cloning site and ribosome binding site of which the ATG initiation codon is located at position 232-234 267-772 : rrnBT₁T₂ containing HindIII-SspI fragment from pKK223 (Pharmacia) 773-3300 : tetracycline resistance gene and origin replication containing EcoRI-DraI fragment of pAT 153 (Bioexcellence)

Table 7 corresponds to the complete restriction site analysis of pIG2.

3169

312

2774

Table 7

Total number of bases is: 3301.
Analysis done on the complete sequence.

List of cuts by enzyme.

				3285				2499														2240	
				3270				2411						3084								1857	3129
				2614				2162						2692								1502	2765
				2211				2038		2499				1751	2512			2930				1381	2041
	2882			1589				1859		2411				1748	1879			696	2917			1368	1815
	2864			1363				1817		2162			2888	1635	1855			883	2195			632	1150
	2750			1227				595		1859			2867	1617	1108			785	881	2808		467	298
2647	2093			970				505		1817		2922	2753	1198	1105	2567	2369	247	784	807	2243	357	265
252	617	1527	222	268	1118	1213	208	376	1872	376	1855	239	2096	271	899	1704	2135	15	234	737	264	213	4
••	••	••	••	••	••	••	••	••	••	••	••	••	••	••	••	••	••	••	••	••	••	••	••
н			III			Ľ		H	Н	II	н	HI	Н		* H	HI.	H					IN	
Acc	Acy	Afl	Aha	Alu	Alw	Apa	Asp	Asu	Ava	Ava	Bal	Bam	Bbe	Bbv	Bbv	Bbv	Bgl	Bin	Bin	Bsp	Bsp	Bst	Cau

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																																		_		
			97	150	204	247	300				217											,	162	227		977								225		
			673	α	03	42	96				2156									,	3167	(1606	19		21195							3069	2039	3127	
			657	1461	01	38	90				1839										2238	1	1603	2136	3073	7000						2381	2888	1855	3004	
			596	1363	97	2360	82				963									1	1855	,	1485	2091	3002	2026						2309	2867	_; ,	2900	
			0	1227	89	ぜ	81			1717	888									1	1500	,	1330	2012	2999	1911						2252	2807	1505	2776	
Con't)	3137		ထ	1222	85	┙	77			1253	877										1379		1187	1893	2721	ლ ⊾	7 324					1855	2753	1487	2768	2658
Table 7 (Con't)	2896	3002	303	1197	81	2309	9/			844	869	2951									1366		1122	1890	2718	1763	2598					1516	2526	u)	2704	9
Te	2887	&	273	1118	1808	2252	2704	3285		546	791	2924				3067			2886		633		1119	1883	2684	1666	2483		3151			1505	2096	673	2470	1917
	2528	9/	268	1053	9	2211	68	7		490		2833	2774	2769		2805			2865		465		913	1869	2577	1484	2354	3199	2297			1053	1657	657	2381	625
	2368	35	9	1042	58	19	63	S	13	Ō	241	~	S	1850		2524			2751	2674	355		361	1740	2526	90	2327	9	2252			673	1287	296	2360	181
	01	1853	19	9	51	8	61	12	0	3	6	\sim	1817	~	444	1655	2	985	2094	196	\sim	3114	260	~	2361	(2322	297	645	Q	2471	657	423	п,	2309	158
	••	•• ••	• ••						••	••	••		••	••	••	 H	••	••	••	••	••	••	••			••		••	••	••	••	••	••	••		••
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	Cfr	Cfr	Cvi						Cvi	Dde	Dpn		Dra	Dsa	Eco	Eco	Eco	Eco	Eco	Eco	Eco	ECO	Fnu			Fnu		Fok	Fok	Gsn	Gsu	Hae	Hae	Hae		Hga

					188	304		8	303	•				227	i I							216	1					291	1							210	1
					1846	2887		1844	2885)				2041		3184						2154	• • • •					2693) !							2051	4) }
			3221		1656	2866		1654	2864	0			2666	2015	3138	2902						1837	1					2426								1844	3297
			3178		1623	2806		1621	2804				2446	1814	3129	2857					3172	961	l)					2124		3265						1710	2950
			2884		1353	2752		1351	2750				2292	1322	2912	2003					3084	886	•	2572				2063		3243						1531	2812
con't)		3025	2863		1286	2600		1284	2598				1994	1175	2897	1782				2398	2416	875	•	2301			2512	2025		823						811	2739
7 (001	2897	2714	2749		1186	2525		18	2523	1			1773	1149	2888	961			3069	2342	2149	867		1709			2328	1998		646						741	2593
Table	2343	2127	2531		1012	2485		1010	2483) - -			1628	959	2765	545			1810	1753	1170	789	2922	1661		3081	1830	1632		315		2898	2885			449	2554
	2311	1836	2092		903	2354		901	2352	;	2648		1553	598	2605	181			1034	1729	1107	777	2831	799		2994	1424	1349		221	3039	2530	2864			394	2368
	1415	1217	2008	2830	422	2095	3200	420	2093	3198	253	3268	1157	564	2529	138			781	826	991	239	2472	304	2826	1318	1100	593	3176	186	1943	2370	2750			234	2251
	837	139	208	2816	371	1944	3068	369	1942	3066	107	766	249	m	2369	94	9	212	246	580	169	7	2200	207	870	1134	253	208	3116	179	1845	2016	2093	230	3068	166	2123
	••	••	••	••	••			••			••	••	••	••		••	••	••	••	••	••	••		••	••	••	••	••		••	••	••	••	••	••	••	
	*I	AI	CI	JII	 			PlI			Η		H	\mathbf{H}		H	*H	н	H	H		Н		H		H	н	* H			H					III	
	Hga	Hgi	Hgi	Hgi	Hha			Hin			Hind	Hin	Hin	Hpa		Hph	Hph	Kpn	Mae	Mae	Mae	Mbo		Mbo	Mbo	Mme	Mnl	Mnl		Mse	Mst	Nae	Nar	Nco	Nhe	Nla	

	σ.											_	>						c	>														
	209											150	7						150	7														
	2045 3223											1281	TOCT		,	3187			1270	13/3														
	2010 3180											1260	1200		,	3168			1366	1.300														
	1975 2970											1160	OCTT	6	3025	2769			•	1148			2275	C/7C										
	1861 2924											7.07	63.5	3169	2830	2763			0	633	3167			7960					0	7767				
<u></u>	1818 2886											i	298	3129	2816	2130	3173	3084	1	296	3127		1700	7 697					•	988				
7 (Con't)	1499 2865											1	265	2765	2714	1928	3060	2895	,	563	2763			2112					1	875				
Table 7	1460 2775											•	467	2240	2127	1850	2649	2883		465	2238			2031						789			<u>,</u>	•
	378 2751		2160					1	2777					~		1367	~	~		355	2039		•	1429				943		777			689	
	241 2533		1189	2739	1983	1636	2660	1859	1862		3136	2646	213	1857	1217	230	647	1867		211	1855		1928	495			936	904		239			Cute is	
	210 2412	\sim	Ť	S	93	25	15	$\boldsymbol{\vdash}$	82	ဖ	_	വ	4	_	13			43	(1.)		_	226	CT)	шı		ניו	1.1	515	4		u ,	296	ک در در در در در در در در در در در در در	70
	••	••	••	••	••	••	••	••	••	••	••	••	••		••	••	••	••	••	**		••	••	••	••	••	••	*	••	••	••	•• .	. 64	ragumu
	a IV	u I		p HI								T T						*IN e				I di	ı y	ığ.	iq IIB	nd IIB*	:h11111		la I	II ot	na III	Xmn I	1 MO.	otal nu
	Nla	Nrn	NS	NS	Pf	Pl	Pl	Ppu	Ps	Ps	Ra	Sa	S)	S.	S	ţ.	Sfa	ST	S		S	St	Ta	Ħ	Ţ	Ŧ	It	×	×	×	×.	בֿ בֿ).T.

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Table 7 (con't)

List of non cutting selected enzymes.

Aat II , Afl II , Apa I , Asu II , Avr II Bgl II , Bsp MI , Bsp MI , Bst EII Eco 311* , Esp I , Hpa I , Mlu I , Mme I , Sca I , Sci I , Sfi I , Sma I , Sho I , Xma I

Bcl Dra Not Sac Spe Xca

> Nde I Sac I Sna BI Vsp I

, Bbv II*

Total number of selected enzymes which do not cut: 44

Figure 13 corresponds to the amino acid sequence of the total fusion protein mTNF-His $_6$ -P $_{32}$.

On this figure:

- the continuous underlined sequence (_____)
 represents the mTNF sequence (first 25 amino acids),
- the dotted underlined sequence (----) represents the polylinker sequence,
- the double underlined sequence (-----) represents the extra amino acids created at cloning site, and
- the amino acid marked with nothing is the antigen sequence starting from the amino acid at position 4 of figure 5.

Figure 14a and 14b correspond to the expression of the mTNF-His $_6$ -P $_{32}$ fusion protein in K12 $_{\Lambda}$ H, given in Example VI, with Fig. 14a representing the Coomassie Brilliant Blue stained SDS-PAGE and 14b representing immunoblots of the gel with anti-32-kDa and anti-mTNF-antibody.

On fig. 14a, the lanes correspond to the following:

Lanes

 protein molecular weight marke 	1.	protein	molecular	weight	markers
--	----	---------	-----------	--------	---------

2.	pmTNF-MPH-Mt32	28°C	1 h	induction
3.	11	42°C		11
4.	**	42°C	2 h	induction
5.	11	42°C	3 h	11
6.	Ħ	28°C	4 h	**
7.	41	42°C	4 h	**
8.	τι	28°C	5 h	**
9.	11	42°C	5 h	**

On fig. 14b, the lanes correspond to the following:

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Lane	2 S				
1.	pmTNF-MPH-Mt32	28°C	1	h	induction
2.	11	42°C	1	h	88
3.	II .	28°C	4	h	**
А	11	42°C	4	h	11

Figure 15 corresponds to the IMAC elution profile of the recombinant antigen with decreasing pH as presented in Example VII.

Figure 16 corresponds to the IMAC elution profile of the recombinant antigen with increasing imidazole concentrations as presented in Example VII.

Figure 17 corresponds to the IMAC elution profile of the recombinant antigen with a step gradient of increasing imidazole concentrations as presented in Example VII.

EXAMPLE I:

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MATERIAL AND METHODS

Screening of the Agtll M. tuberculosis recombinant DNA library with anti-32-kDa antiserum

Agt11 recombinant library constructed genomic DNA of M. tuberculosis (Erdman strain), was obtained from R. Young (35). Screening was performed as described (14,35) with some modifications hereafter mentioned. Agt11 infected E. coli Y1090 (105 pfu per 150 mm plate) were seeded on NZYM plates (Gibco)(16) and incubated at 42°C for 24 hrs. To induce expression of the β -galactosidase-fusion proteins the plates were overlaid with isopropyl β -D-thiogalactoside (IPTG)saturated filters (Hybond C extra, Amersham), incubated for 2 hrs at 37°C. Screening was done with a antiserum. anti-32-kDa polyclonal rabbit polyclonal antiserum rabbit anti-32-kDa antiserum was obtained by raising antiserum against the P32 M. bovis BCG (strain 1173P2 - Institut Pasteur Paris) follows: 400 μ g (purified P₃₂ protein of M. bovis BCG) per ml physiological saline were mixed with one volume of incomplete Freund's adjuvant. The material was homogenized and injected intradermally in 50 μ l doses, delivered at 10 sites in the back of the rabbits, at 0, 4, 7 and 8 weeks (adjuvant was replaced by the diluent for the last injection). One week later, the rabbits were bled and the sera tested for antibody level before being distributed in aliquots and stored at -80°C.

The polyclonal rabbit anti-32-kDa antiserum was pre-absorbed on E. coli lysate (14) and used at a final dilution of 1:300. A secondary alkaline-phosphatase anti-rabbit IgG conjugate (Promega), diluted at 1:5000 was used to detect the β -galactosidase fusion proteins. For color development nitro blue tetrazolium (NBT) and 5-bromo-4-chloro-3-indolyl phosphate (BCIP) were used. Reactive areas on the filter turned deep purple within 30 min. Usually three consecutive purification steps were performed to obtain pure clones. IPTG, BCIP and NBT were from Promega corp. (Madison WI.).

Plaque screening by hybridization for obtaining the secondary clones BY1, By2 and By5 hereafter defined

The procedure used was as described by Maniatis et al. (14).

Preparation of crude lysates from \(\lambda\)gtll recombinant lysogens

Colonies of E. coli Y1089 were lysogenized with appropriate λ gtl1 recombinants as described by Hyunh et al. (14). Single colonies of lysogenized E. coli Y1089 were inoculated into LB medium and grown to an optical density of 0.5 at 600nm at 30°C. After a heat shock at 45°C for 20 min., the production of β -galactosidase-fusion protein was induced by the addition of IPTG to a final concentration of 10 mM. Incubation was continued for 60 min. at 37°C and cells were quickly harvested by centrifugation. Cells were concentrated 50 times in buffer (10 mM Tris pH 8.0, 2 mM EDTA) and rapidly frozen into liquid nitrogen. The samples were lysed by

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thawing and treated with 100 μ g/ml DNase I in EcoRI restriction buffer, for 5-10 minutes at 37°C. Immunoblotting (Western blotting) analysis:

electrophoresis, recombinant SDS-PAGE After lysogen proteins were blotted onto nitrocellulose membranes (Hybond C, Amersham) as described by Towbin et al. (29). The expression of mycobacterial antigens, coli in E. Y1089 fused to eta-galactosidase visualized by the binding of a polyclonal rabbit anti-32-kDa antiserum (1:1000) obtained as described in above paragraph "Screening of the Agt11 Μ. tuberculosis recombinant DNA library with anti-32-kDa antiserum" and using a monoclonal anti- β -galactosidase antibody (Promega). A secondary alkaline-phosphatase anti-rabbit IgG conjugate (Promega) diluted at 1:5000, was used to detect the fusion proteins.

The use of these various antibodies enables to detect the β -galactosidase fusion protein. This reaction is due to the <u>M. tuberculosis</u> protein because of the fact that non fused- β -galactosidase is also present on the same gel and is not recognized by the serum from tuberculous patients.

In order to identify selective recognition of recombinant fusion proteins by human tuberculous sera, nitrocellulose sheets were incubated overnight with these sera (1:50)(after blocking aspecific protein human tuberculous sera were sites). The selected for their reactivity (high or low) against the purified 32-kDa antigen of M. bovis BCG tested in a Dot blot assay as previously described (31). Reactive areas nitrocellulose sheets were revealed incubation with peroxidase conjugated goat anti-human IgG antibody (Dakopatts, Copenhagen, Denmark) (1:200) for 4 hrs and after repeated washings color reaction was developed by adding peroxidase substrate (α - chloronaphtol) (Bio-Rad) in the presence of peroxidase and hydrogen peroxide.

Recombinant DNA analysis

identification of M. tuberculosis inserts in purified Agt11 clones was performed by EcoRI restriction. After digestion, the excised inserts were run on agarose gels and submitted to Southern hybridization. Probes were labeled with α^{32} P-dCTP by random priming (10). Other restriction sites were located by single and double digestions of recombinant Agt11 phage DNA or their subcloned EcoRI fragments by HindIII, PstI, KpnI, AccI and SphI.

Sequencing

Sequence analysis was done by the primer extension dideoxy termination method of Sanger et al. (25) after subcloning of specific fragments in Bluescribe-M13 (6) or in mp10 and mp11 M13 vectors (Methods in Enzymology, 1983, p.20-89, Joachim Messing, vectors for cloning, Academic Press). Sequence analysis was greatly hampered by the high GC content of the M. tuberculosis DNA (65%). Sequencing reactions were therefore performed with several DNA polymerases: T7 DNA polymerase ("Sequenase" USB), Klenow fragment of DNA polymerase I (Amersham) and in some cases with AMV reverse transcriptase (Super RT, Anglian Biotechnology Ltd.) and sometimes with dITP instead of dGTP. Several oligodeoxynucleotides were synthesized and used to focus ambiguous regions of the sequence. The sequencing strategy is summarized in Fig. 2 In order to trace possible artefactual frameshifts in some ambiguous regions, a special program was used to define the most probable open reading frame in sequences containing a high proportion of GC (3). Several regions particularly to sequencing artefacts prone were confirmed chemical corrected by sequencing (18).purpose, fragments were subcloned in the chemical

sequencing vector pGV462 (21) and analysed as described previously. Selected restriction fragments of about 250-350bp were isolated, made blunt-ended by treatment with either Klenow polymerase or Mung bean nuclease, and subcloned in the SmaI or HincII site of pGV462. Both strands of the inserted DNA were sequenced by single-end labeling at Tth 111I or BstEII (32) and a modified chemical degradation strategy (33).

Routine computer aided analysis of the nucleic acid and deduced amino acid sequences were performed with the LGBC program from Bellon (2). Homology searches used the FASTA programs from Pearson and Lipman (23) and the Protein Identification Resource (PIR) from the National Biomedical Research Fundation - Washington (NBRF) (NBRF/PIR data bank), release 16 (march 1988).

RESULTS

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- Screening of the AgtllM, M. tuberculosis recombinant DNA library with polyclonal anti-32-kDa antiserum:

Ten filters representing 1.5x10⁶ plaques were probed with a polyclonal rabbit anti-32-kDa antiserum (8). Following purification, six independent positive clones were obtained.

Analysis of recombinant clones

EcoRI restriction analysis of these 6 purified λgtl1 recombinant clones DNA, (Fig. 1A) revealed 4 different types of insert. Clone 15 had an insert with a total length of 3.8 kb with two additional internal EcoRI sites resulting in three DNA fragments of 1.8 kb, 1.5 kb and 0.5 kb. The DNA Insert of clone 16 was 1.7 kb long. Clones 17 and 19 had a DNA insert of almost identical length being 2.7 kb and

2.8 kb respectively.

Finally, clone 23 (not shown) and clone 24 both contained an insert of 4 kb with one additional EcoRI restriction site giving two fragments of 2.3 kb and

1.7 kb. Southern analysis (data not shown) showed that the DNA inserts of clones 15, 16, 19 and the small fragment (1.7 kb) of clone 24 only hybridized with themselves whereas clone 17 (2.7 kb) hybridized with itself but also equally well with the 2.3 kb fragment of clone 24. Clones 15, 16 and 19 are thus distinct and unrelated to the 17, 23, 24 group. This interpretation was further confirmed by analysis of crude lysates of E. coli Y1089 lysogenized with the appropriate Agt11 recombinants and induced with IPTG. Western blot analysis (Fig. 1B), showed no fusion protein, either mature or incomplete, reactive with the polyclonal anti-32-kDa antiserum in cells expressing clones 15, 16 and 19. Clones 15, 16 and 19, were thus considered as false positives and were not further studied. On the contrary, cells lysogenized with clone 23 and 24 produced an immunoreactive fusion protein containing about 10 kDa of the 32-kDa protein. Clone 17 finally expressed a fusion protein of which the foreign polypeptide part is about 25 kDa long. The restriction endonuclease maps of the 2.3 kb insert of clone 24 and of the 2.7 kb fragment of clone 17 (Fig. 2) allowed us to align and orient the two inserts suggesting that the latter corresponds to a ±0.5 kb 5' extension of the first.

clone 17 incomplete, the As was recombinant M. tuberculosis DNA library was screened by hybridization with a 120 bp EcoRI-Kpnl restriction fragment corresponding to the very 5' end of the DNA insert of clone 17 (previously subcloned in a Blue Scribe vector commercialized by Vector cloning Systems (Stratagene Cloning System) (Fig.2). Three 5'-extended clones By1, By2 and By5 were isolated, analyzed by restriction and aligned. The largest insert, contained the information for the entire coding region (see below) flanked by 3.1 kb upstream and 1.1 kb downstream (Fig. 2).

DNA sequencing

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The 1358 base pairs nucleotide sequence derived various λgt11 overlapping represented in Fig. 3a and Fig. 3b. The DNA sequence contains a 1059 base pair open reading frame starting at position 183 and ending with a TAG codon at position occurs that the NH2-terminal amino-acid sequence, (phe-ser-arg-pro-gly-leu-pro-valglu-tyr-leu-gln-val-pro-ser-pro-ser-met-gly-arg-aspile-lys-val-gln-phe-gln-ser-gly-gly-ala-asn) which can be located within this open reading frame from the nucleotide sequence beginning with a TTT codon at position 360 corresponds to the same NH2-terminal amino acid sequence of the MPB 59 antigen except for the amino acids at position 20, 21, 31, which respectively gly, cys and asn in the MPB 59 (34). Therefore, the DNA region upstream of this sequence is expected to encode a signal peptide required for the secretion of a protein of 32-kDa. The mature protein thus presumably consists of 295 amino acid residues from the N-terminal Phe (TTT codon) to the C-terminal Ala (GCC codon) (Fig. 5).

Six ATG codons were found to precede the TTT at position 360 in the same reading frame. Usage of any of these ATGs in the same reading frame would lead to the synthesis of signal peptides of 29,42,47,49,55 and 59 residues.

Hydropathy pattern

The hydropathy pattern coding sequence of the protein of 32-kDa of the invention and that of the antigen α of BCG (17) were determined by the method of Kyte and Doolittle (15). The nonapeptide profiles are shown in Fig. 6. Besides the initial hydrophobic signal peptide region, several hydrophilic domains could be

identified. It is interesting to note that the overall hydrophilicity pattern of the protein of 32-kDa of the invention is comparable to that of the BCG antigen α . For both proteins, a domain of highest hydrophilicity could be identified between amino acid residues 200 and 250.

Homology

Matsuo et al. (17) recently published the sequence of a 1095 nucleotide cloned DNA corresponding to the gene coding for the antigen α of BCG. The 978 bp coding region of M. bovis antigen α as revised in Infection and Immunity, vol. 58, p. 550-556, 1990, and 1017 bp coding regions of the protein of 32-kDa of invention show a 77.5% homology, in an aligned region of 942 bp. At the amino acid level both precursor protein sequences share 75.6% identical residues. addition, 17.6% of the amino acids correspond evolutionary conserved replacements as defined in the algorithm used for the comparison (PAM250 matrix, ref 23). Figure 7 shows sequence divergences in the Nterminal of the signal peptide. The amino terminal sequence - 32 amino acids - of both mature proteins is identical except for position 31.

Human sera recognize the recombinant 32-kDa protein

Fig. 8 shows that serum samples from tuberculous patients when immunoblotted with a crude <u>E. coli</u> extract expressing clone 17 distinctly react with the 140 kDa fusion protein (lanes 4 to 6) contain the protein of 32-kDa of the invention, but not with unfused β -galactosidase expressed in a parallel extract (lanes 10 to 12). Serum samples from two negative controls selected as responding very little to the purified protein of 32-kDa of the invention does neither recognize the 140 kDa fused protein containing the protein of 32-kDa of the invention, nor the unfused β -galactosidase (lanes 2, 3 and 8 and 9). The 140 k-Da

fused protein and the unfused β -galactosidase were easily localized reacting with the anti- β -galactosidase monoclonal antibody (lanes 1 to 7).

The invention has enabled to prepare a DNA region coding particularly for a protein of 32-kDa fig.5); said DNA region containing an open reading frame of 338 codons (stop codon non included). At position 220 a TTT encoding the first amino acid of the mature protein is followed by the 295 triplets coding for the mature protein of 32-kDa. The size of this open reading frame, the immunoreactivity of the derived fusion proteins, the presence of a signal peptide and, especially, the identification within this gene of a NH2-terminal region highly homologous to that found in the MPB 59 antigen (31/32 amino acids homology) and in the BCG antigen α (31/32 amino acids homology) (see Fig. 7), strongly suggest that the DNA fragment described contains the complete cistron encoding the protein of 32-kDa secreted by M. tuberculosis, and which had never been so far identified in a non ambiguous way.

Six ATG codons were found to precede this TTT at position 220 in the same reading frame. Usage of any of these ATGs in the same reading frame would lead to the synthesis of signal peptides of 43, 48, 50, 56 or 60 residues. Among these various possibilities, initiation is more likely to take place either at ATG₉₁ or ATG₅₂ because both are preceded by a plausible <u>E. coli</u>-like promoter and a Shine-Dalgarno motif.

If initiation takes place at ATG₉₁, the corresponding signal peptide would code for a rather long peptide signal of 43 residues. This length however is not uncommon among secreted proteins from Gram positive bacteria (5). It would be preceded by a typical <u>E. coli</u> Shine-Dalgarno motif (4/6 residues homologous to AGGAGG) at a suitable distance.

If initiation takes place at ATG_{52} , the corresponding signal peptide would code for a peptide signal of 56 residues but would have a less stringent Shine-Dalgarno ribosome binding site sequence.

The region encompassing the translation termination triplet was particularly sensitive to secondary structure effects which lead to so-called compressions on the sequencing gels. In front of the TAG termination codon at position 1105, 22 out of 23 residues are G-C base pairs, of which 9 are G's.

Upstream ATG130, a sequence resembling an E. coli promoter (11) comprising an hexanucleotide (TTGAGA) (homology 5/6 to TTGACA) and a AAGAAT box (homology 4/6 to TATAAT) separated by 16 nucleotides was observed. Upstream the potential initiating codon ATGo1, could detect several sequences homologous to the E. coli "-35 hexanucleotide box", followed by a sequence TATAAT box. Among these, resembling a the suggestive is illustrated on Fig. 3a and 3b. It comprises a TTGGCC at position 59 (fig. 3a 3b) (homology 4/6 to TTGACA) separated by 14 nucleotides from a GATAAG (homology 4/6 to TATAAT). Interestingly this putative promoter region shares no extensive sequence homology with the promoter region described for the BCG protein α -gene (17) nor with that described for the 65 kDa protein gene (26, 28).

Searching the NBRF data bank (issue 16.0) any significant homology between the protein of 32-kDa of the invention and any other completely known protein sequence could not be detected. In particular no significant homology was observed between the 32-kDa protein and α and β subunits of the human fibronectin receptor (1). The NH₂-terminal sequence of the 32-kDa protein of the invention is highly homologous - 29/32 amino acids - to that previously published for BCG MPB 59 antigen (34) and to that of BCG α -antigen - 31/32

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amino acids - (Matsuo, 17) and is identical in its first 6 amino acids with the 32-kDa protein of \underline{M} . bovis BCG (9). However, the presumed initiating methionine precedes an additional 29 or 42 amino acid hydrophobic sequence which differs from the one of α -antigen (cf. Fig. 7), but displaying all the characteristics attributed to signal sequences of secreted polypeptides in prokaryotes (22).

Interestingly, no significant homology between the nucleic acid (1-1358) of the invention (cf. fig. 3a and 3b) and the DNA of the antigen α of Matsuo exists within their putative promoter regions.

EXAMPLE II: CONSTRUCTION OF A BACTERIAL PLASMID CONTAINING THE ENTIRE CODING SEQUENCE OF THE 32-kDa PROTEIN OF M. TUBERCULOSIS

In the previous example, in figure 2, the various overlapping λ gtll isolates covering the 32-kDa protein gene region from M. tuberculosis were described. Several DNA fragments were subcloned from these λ gtll phages in the Blue Scribe M13+ plasmid (Stratagene). Since none of these plasmids contained the entire coding sequence of the

32-kDa protein gene, a plasmid containing this sequence was reconstructed.

Step 1 : Preparation of the DNA fragments :

- 1) The plasmid BS-By5-800 obtained by subcloning HindIII fragments of By5 (cf. fig. 2) into the Blue Scribe M13⁺ plasmid (Stratagene), was digested with HindIII and a fragment of 800 bp was obtained and isolated from a 1% agarose gel by electroelution.
- 2) The plasmid BS-4.1 obtained by subcloning the 2,7 kb EcoRI insert from λgt11-17) into the Blue Scribe M13⁺ plasmid (Stratagene) (see fig.2 of patent application) was digested with HindIII and SphI and a fragment of 1500 bp was obtained and isolated from a 1% agarose gel by electroelution.

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3) Blue Scribe M13* was digested with HindIII and SphI, and treated with calf intestine alkaline phosphatase (special quality for molecular biology, Boehringer Mannheim) as indicated by the manufacturer.

Step 2 : ligation :

The ligation reaction contained:

125 ng of the 800 bp HindIII fragment (1)

125 ng of the 1500 bp SphI-HindIII insert (2)

50 ng of the HindIII-SphI digested BSM13* vector (3)

2 μ l of 10 ligation buffer (Maniatis et al., 1982)

1 μ l of (= 2,5 U) of T4 DNA ligase (Amersham)

4 μ l PEG 6000, 25% (w/v)

8 µ1 H₂O

The incubation was for 4 hours at 16°C.

Step 3 : Transformation :

100 μ l of DH5 α <u>E. coli</u> (Gibco BRL) were transformed with 10 μ l of the ligation reaction (step 2) and plated on IPTG, X-Gal ampicillin plates, as indicated by the manufacturer. About 70 white colonies were obtained.

step 4:

As the 800 bp fragment could have been inserted in both orientations, plasmidic DNA from several clones were analyzed by digestion with PstI in order to select one clone (different from clone 11), characterized by the presence of 2 small fragments of 229 and 294 bp. This construction contains the HindIII-HindIII-SphI the correct complex in orientation. The plasmid containing this new construction was called "BS.BK.P32.complet".

EXAMPLE III: EXPRESSION OF A POLYPEPTIDE OF THE INVENTION IN E. COLI:

The DNA sequence coding for a polypeptide, or part of it, can be linked to a ribosome binding site which is part of the expression vector, or can be fused to }

the information of another protein or peptide already present on the expression vector.

In the former case the information is expressed as such and hence devoid of any foreign sequences (except maybe for the aminoterminal methionine which is not always removed by \underline{E} . \underline{coli}).

In the latter case the expressed protein is a hybrid or a fusion protein.

The gene, coding for the polypeptide, and the expression vector are treated with the appropriate restriction enzyme(s) or manipulated otherwise as to The resulting allowing ligation. termini recombinant vector is used to transform a host. The transformants are analyzed for the presence and proper orientation of the inserted gene. In addition, the cloning vector may be used to transform other strains of a chosen host. Various methods and materials for preparing recombinant vectors, transforming them to host cells and expressing polypeptides and proteins are described by Panayatatos, N., in "Plasmids, a practical approach (ed. K.G. Hardy, IRL Press) pp.163-176, by Old and Primrose, principals of gene manipulation (2d Ed, 1981) and are well known by those skilled in the art.

Various cloning vectors may be utilized for expression. Although a plasmid is preferable, the vector may be a bacteriophage or cosmid. The vector chosen should be compatible with the host cell chosen.

Moreover, the plasmid should have a phenotypic property that will enable the transformed host cells to be readily identified and separated from those which are not transformed. Such selection genes can be a gene providing resistance to an antibiotic like for instance, tetracyclin, carbenicillin, kanamycin, chloramphenicol, streptomycin, etc.

In order to express the coding sequence of a gene in \underline{E} . \underline{coli} the expression vector should also contain

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the necessary signals for transcription and translation.

Hence it should contain a promoter, synthetic or derived from a natural source, which is functional in <u>E. coli</u>. Preferably, although usually not absolutely necessary, the promoter should be controllable by the manipulator. Examples of widely used controllable promoters for expression in <u>E. coli</u> are the lac, the trp, the tac and the lambda PL and PR promoter.

Preferably, the expression vector should also contain a terminator of transcription functional in \underline{E} . \underline{coli} . Examples of used terminators of transcription are the trp and the rrnB terminators.

Furthermore, the expression vector should contain a ribosome binding site, synthetic or from a natural source, allowing translation and hence expression of a downstream coding sequence. Moreover, when expression devoid of foreign sequences is desired, a unique restriction site, positioned in such a way that it allows ligation of the sequence directly to the initiation codon of the ribosome binding site, should be present.

A suitable plasmid for performing this type of expression is pKK233-2 (Pharmacia). This plasmid contains the trc promoter, the lac Z ribosome binding site and the rrnB transcription terminator.

Also suitable is plasmid pIGRI (Innogenetics, Ghent, Belgium). This plasmid contains the tetracycline resistance gene and the origin of replication of pAT₁₅₃ (available from Bioexcellence, Biores B.V., Woerden, The Netherlands), the lambda PL promoter up to the MboII site in the 5' untranslated region of the lambda N gene (originating from pPL(λ); Pharmacia).

Downstream from the PL promoter, a synthetic sequence was introduced which encodes a "two cistron" translation casette whereby the stop codon of the first

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cistron (being the first 25 amino acids of TNF, except for Leu at position 1 which is converted to Val) is situated between the Shine-Dalgarno sequence and the initiation codon of the second ribosome binding site. The restriction and genetic map of pIGRI is represented in Fig. 10a.

Fig. 10b and Table 5 represent respectively the nucleic acid sequence and complete restriction site analysis of pIGRI.

However, when expression as a hybrid protein is desired, then the expression vector should also contain the coding sequence of a peptide or polypeptide which is (preferably highly) expressed by this vector in the appropriate host.

In this case the expression vector should contain a unique cleavage site for one or more restriction endonucleases downstream of the coding sequence.

Plasmids pEX1, 2 and 3 (Boehringer, Mannheim) and pUEX1, 2 and 2 (Amersham) are useful for this purpose.

They contain an ampicillin resistance gene and the origin of replication of pBR322 (Bolivar at al. (1977) Gene 2, 95-113), the lac Z gene fused at its 5' end to the lambda PR promoter together with the coding sequence for the 9 first amino acids of its natural gene cro, and a multiple cloning site at the 3' end of the lac Z coding sequence allowing production of a beta galactosidase fused polypeptide.

The pUEX vectors also contain the CI857 allele of the bacteriophage lambda CI repressor gene.

Also useful is plasmid pmTNF MPH (Innogenetics). It contains the tetracycline resistance gene and the origin of replication of pAT₁₅₃ (obtainable from Bioexcellence, Biores B.V., Woerden. The Netherlands), the lambda PL promoter up to the MboII site in the N gene 5' untranslated region (originating from pPL(λ); Pharmacia), followed by a synthetic ribosome binding

site (see sequence data), and the information encoding the first 25 AA of mTNF (except for the initial Leu which is converted to Val). This sequence is, in turn, followed by a synthetic polylinker sequence which encodes six consecutive histidines followed by several proteolytic sites (a formic acid, CNBr, kallikrein, and E. coli protease VII sensitive site, respectively), each accessible via a different restriction enzyme which is unique for the plasmid (SmaI, NcoI, BspMII and StuI, respectively; see restriction and genetic map, Fig. 11a). Downstream from the polylinker, several transcription terminators are present including the E. trp terminator (synthetic) and the rrnBT₁T₂ (originating from pKK223-3; Pharmacia). The nucleic acid sequence of this plasmid is represented in Fig. 11b.

Table 6 gives a complete restriction site analysis of pmTNF MPH.

The presence of 6 successive histidines allows purification of the fusion protein by Immobilized Metal Ion Affinity Chromatography (IMAC).

After purification, the foreign part of the hybrid protein can be removed by a suitable protein cleavage method and the cleaved product can then be separated from the uncleaved molecules using the same IMAC based purification procedure.

In all the above-mentioned plasmids where the lambda PL or PR promoter is used, the promoter is temperature-controlled by means of the expression of the lambda cI ts 857 allele which is either present on a defective prophage incorporated in the chromosome of the host (K12AH, ATCC n° 33767) or on a second compatible plasmid (pACYC derivative). Only in the pUEX vectors is this cI allele present on the vector itself.

It is to be understood that the plasmids presented above are exemplary and other plasmids or types of

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expression vectors maybe employed without departing from the spirit or scope of the present invention.

If a bacteriophage or phagemid is used, instead of plasmid, it should have substantially the same characteristics used to select a plasmid as described above.

EXAMPLE IV : SUBCLONING OF THE P32 ANTIGEN IN PLASMID PIGRI :

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Fifteen μ g of plasmid "BS-BK-P₃₂ complet" (see Example II) was digested with <u>EclXI</u> and <u>BstEII</u> (Boehringer, Mannheim) according to the conditions recommended by the supplier except that at least 3 units of enzyme were used per μ g of DNA. <u>EclXI</u> cuts at position 226 (Fig. 5) and <u>BstEII</u> at position 1136, thus approaching very closely the start and stop codon of the mature P₃₂ antigen. This DNA is hereafter called DNA coding for the "P₃₂ antigen fragment".

The DNA coding for the " P_{32} antigen fragment" (as defined above) is subcloned in pIGRI (see fig. 10a) for expression of a polypeptide devoid of any foreign sequences. To bring the ATG codon of the expression vector in frame with the P_{32} reading frame, an intermediary construct is made in pIG2 (for restriction and genetic map, see fig. 12a; DNA sequences, see fig. 12b; complete restriction site analysis, see Table 7).

Five μg of plasmid pIG2 is digested with NCoI. Its 5' sticky ends are filled in prior to dephosphorylation.

Therefore, the DNA was incubated in 40 μ l NB buffer (0.05 M Tris-Cl pH 7.4; 10 mM MgCl₂; 0.05% β -mercaptoethanol) containing 0.5 mM of all four dXTP (X = A,T,C,G) and 2 μ l of Klenow fragment of <u>E. coli</u> DNA polymerase I (5 U/ μ l, Boehringer, Mannheim) for at least 3 h at 15°C.

After blunting, the DNA was once extracted with one volume of phenol equilibrated against 200 mM Tris-

Cl 8, twice with Hq at least two volumes diethylether and finally collected using the "gene clean kit .M. (Bio101) as recommended by the supplier. The DNA was then dephosphorylated at the 5' ends in 30 μl of CIP buffer (50 mM TrisCl pH 8, 1 mM ZnCl₂) and 20 units of calf intestine phosphatase (high concentration, Boehringer, Mannheim). The mixture was incubated at 37°C for 30 min, then EGTA (ethyleneglycol bis $(\beta$ -aminoethylether)-N,N,N',N' tetraacetic acid) pH 8 is added to a final concentration of 10 mM. The mixture was then extracted with phenol followed by diethylether as described above, and the DNA was precipitated by addition of 1/10 volume of 3 M KAc (Ac = CH_3COO) pH 4.8 and 2 volumes of ethanol followed by storage at -20°C for at least one hour.

After centrifugation at 13000 rpm in a Biofuge A (Hereaus) for 5 min the pelleted DNA was dissolved in H_2O to a final concentration of 0.2 $\mu g/\mu l$.

The <u>EclXI-BstEII</u> fragment, coding for the " P_{32} antigen fragment" (see above) was electrophoresed on a 1% agarose gel (BRL) to separate it from the rest of the plasmid and was isolated from the gel by centrifugation over a Millipore HVLP filter (ϕ 2 cm)(2 min,, 13000 rpm, Biofuge at room temperature) and extracted with Tris equilibrated phenol followed by diethylether as described above.

The DNA was subsequently collected using the "Gene clean kit^{T.M.}" (Bio101) as recommended by the supplier.

After that, the 5' sticky ends were blunted by treatment with the Klenow fragment of <u>E. coli</u> DNA polymerase I as described above and the DNA was then again collected using the "Gene clean kit^{I.M.}" in order to dissolve it in 7 μ l of H₂O.

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One μl of vector DNA is added together with one μl of 10 x ligase buffer (0.5 M TrisCl pH 7.4, 100 mM MgCl₂, 5 mM ATP, 50 mM DTT (dithiothreitol)) and 1 μl

of T4 DNA ligase (1 unit/ μ l, Boehringer, Mannheim). Ligation was performed for 6 h at 13°C and 5 μ l of the mixture is then used to transform strain DH1 (lambda) [strain DH1 - ATCC N° 33849 - lysogenized with wild type bacteriophage λ] using standard transformation techniques as described for instance by Maniatis et al. in "Molecular cloning, a laboratory manual", Cold Spring Harbor Laboratory (1982).

Individual transformants are grown and lysed for plasmid DNA preparation using standard procedures (Experiments with gene fusions, Cold Spring Harbor Laboratory (1984) (T.J. Silhavy, H.L. Berman and L.W. Enquist, eds) and the DNA preparations are checked for the correct orientation of the gene within the plasmid by restriction enzyme analysis.

A check for correct blunting is done by verifying the restoration of the $\underline{\text{NcoI}}$ site at the 5' and 3' end of the antigen coding sequence. One of the clones containing the P_{32} antigen fragment in the correct orientation is kept for further work and designated pIG_2 -Mt32. In this intermediary construct, the DNA encoding the antigen is not in frame with the ATG codon. However, it can now be moved as a NcoI fragment to another expression vector.

15 μ g of pIG₂-Mt32 is digested with NcoI. The NcoI fragment encoding the P₃₂ antigen is gel purified and blunted as described above. After purification, using "gene clear kit TM" it is dissolved in 7 μ l of H₂O.

5 μg of plasmid pIGRI is digested with NcoI, blunted and dephosphorylated as described above. After phenol extraction, followed by diethylether and ethanolprecipitation, the pellet is dissolved in H_2O to a final concentration of 0.2 $\mu g/\mu l$.

Ligation of vector and "antigen fragment" DNA is carried out as described above. The ligation mixture is then transformed into strain DH1 (lambda) and

individual transformants are analysed for the correct orientation of the gene within the plasmid by restriction enzyme analysis. A check for correct blunting is done by verifying the creation of a new NsiI site at the 5' and 3' ends of the antigen coding sequence. One of the clones containing the P₃₂ antigen fragment in the correct orientation is kept for further work and designated pIGRI.Mt32.

EXAMPLE V: SUBCLONING OF THE P32 ANTIGEN IN pmTNF MPH:

Fifteen μ g of the plasmid pIG2 Mt32 (see example IV) was digested with the restriction enzyme NCOI (Boehringer, Mannheim), according to the conditions recommended by the supplier except that at least 3 units of enzyme were used per μ g of DNA.

After digestion, the reaction mixture is extracted with phenol equilibrated against 200mM TrisCl pH 8, (one volume), twice with diethylether (2 volumes) and precipitated by addition of 1/10 volume of 3 M KAc (Ac=CH₃COO) pH 4.8 and 2 volumes of ethanol followed by storage at -20°C for at least one hour.

After centrifugation for 5 minutes at 13000 rpm in a Biofuge A (Hereaus) the DNA is electrophoresed on a 1% agarose gel (BRL).

The DNA coding for the " P_{32} antigen fragment" as described above, is isolated by centrifugation over a Millipore HVLP filter (ϕ 2cm)(2 minutes, 13000 rpm, Biofuge at room temperature) and extracted one with trisCl equilibrated phenol and twice with diethylether. The DNA is subsequently collected using "Gene clean kit T.M.M (Bio 101) and dissolved in $7\mu l$ of H_2O .

The 5' overhanging ends of the DNA fragment generated by digestion with NcoI were filled in by incubating the DNA in 40 μ l NB buffer (0.05 M Tris-HCl, pH 7.4; 10 mM MgCl₂; 0.05% β -mercaptoethanol) containing 0.5 mM of all four dXTPS (X = A, T, C, G) and 2μ l of Klenow fragment of E. coli DNA polymerase I

(5 units/ μ l Boehringer Mannheim) for at least 3 h at 15°C. After blunting, the DNA was extracted with phenol, followed by diethylether, and collected using a "gene clean kit T.M." as described above.

Five μg of plasmid pmTNF MPH is digested with StuI, subsequently extracted with phenol, followed by diethylether, and precipitated as described above. The restriction digest is verified by electrophoresis of a 0.5 μg sample on an analytical 1,2% agarose gel.

The plasmid DNA is then desphosphorylated at the 5' ends to prevent self-ligation in $30\mu l$ of CIP buffer (50 mM TrisCl pH 8, 1 mM ZnCl2) and 20 to 25 units of (high concentration, intestine phosphatase calf Boehringer Mannheim). The mixture is incubated at 37°C 30 minutes, then EGTA (ethyleneglycol bis aminoethylether) - N, N, N', N' tetraacetic acid) added to a final concentration of 10 mM. The mixture is extracted with phenol followed by diethylether and the DNA is precipitated as described above. The precipitate is pelleted by centrifugation in a Biofuge A (Hereaus) at 13000 rpm for 10 min at 4°C and the pellet is dissolved in H2O to a final DNA concentration of 0.2 $\mu q/\mu l$.

One μ l of this vector DNA is mixed with the 7 μ l solution containing the DNA fragment coding for the "P32antigen fragment" (as defined above) and 1 μ l 10 x ligase buffer (0.5 M TrisCl pH7.4, 100 mM MgCl2, 5 mM ATP, 50 mM DTT (dithiothreitol)) plus 1 μ l T₄ DNA ligase (1 unit/ μ l, Boehringer Mannheim) is added. The mixture is incubated at 13°C for 6 hours and 5 μ l of the mixture is then used for transformation into strain DH1(lambda) using standard transformation techniques are described by for instance Maniatis et al. in "Molecular cloning, a laboratory manual", Cold Spring Harbor Laboratory (1982).

Individual transformants are grown and then lysed for plasmid DNA preparation using standard procedures

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(Experiments with gene fusions, Cold Spring Harbor Laboratory 1984 (T.J. Silhavy, M.L. Berman and L.W. Enquist eds.)) and are checked for the correct orientation of the gene within the plasmid by restriction enzyme analysis.

One of the clones containing the DNA sequence encoding the antigen fragment in the correct orientation was retained for further work designated pmTNF-MPH-Mt32. It encodes all information of the P_{32} antigen starting from position +4 in the amino acid sequence (see fig. 5). The amino acid sequence of the total fusion protein is represented in fig. 13.

EXAMPLE VI: INDUCTION OF ANTIGEN EXPRESSION FROM pmTNF MPH Mt32:

A- MATERIAL AND METHODS

DNA of pmTNF-MPH-Mt32 is transformed into <u>E. coli</u> strain K12ΔH (ATCC 33767) using standard transformation procedures except that the growth temperature of the cultures is reduced to 28°C and the heat shock temperature to 34°C.

A culture of K12 Δ H harboring pmTNF-MPH-Mt32, grown overnight in Luria broth at 28°C with vigorous shaking in the presence of 10 μ g/ml tetracycline, is inoculated into fresh Luria broth containing tetracyclin (10 μ g/ml) and grown to an optical density at 600 nanometers of 0.2 in the same conditions as for the overnight culture.

When the optical density at 600 nanometers has reached 0.2 half of the culture is shifted to 42°C to induce expression while the other half remains at 28°C as a control. At several time intervals aliquots are taken which are extracted with one volume of phenol equilibrated against M9 salts (0.1% ammonium chloride, 0.3% potassium dihydrogenium phosphate, 1.5% disodium hydrogenium phosphate, 12 molecules of water) and 1% SDS. At the same time, the optical density (600 nm) of

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the culture is checked. The proteins are precipitated from the phenol phase by addition of two volumes of acetone and storage overnight at -20°C. The precipitate is pelleted (Biofuge A, 5 min., 13000 rpm, room temperature) dried at the air, dissolved in a volume of Laemmli (Nature (1970) $\underline{227}$:680) sample buffer (+ β mercapto ethanol) according to the optical density and boiled for 3 min.

Samples are then run on a SDS polyacrylamide gel Laemmli (1970). to according monitored by both is mTNF-His6-P32 induction of (CBB) staining and Blue Brilliant Coomassie immunoblotting. CBB staining is performed by immersing the gel in a 1/10 diluted CBB staining solution (0.5 g CBB-R250 (Serva) in 90 ml methanol : H_2O (1:1 v/v) and 10 ml glacial acetic acid) and left for about one hour on a gently rotating platform. After destaining for a few hours in destaining solution (30% methanol, 7% glacial acetic acid) protein bands are visualised and can be scanned with a densitometer (Ultroscan Enhanced Laser Densitometer, LKB).

For immunoblotting the proteins are blotted onto Hybond C membranes (Amersham) as described by Townbin et al (1979). After blotting, proteins on the membrane are temporarily visualised with Ponceau S (Serva) and the position of the molecular weight markers indicated. The stain is then removed by washing in H_2O . Aspecific protein binding sites blocked are incubating the blots in 10% non-fat dried milk for about 1 hour on a gently rotating platform. After washing twice with NT buffer (25 mM Tris-HCl, pH 8.0; 150 mM NaCl) blots are incubated with polyclonal rabbit anti-32-kDa antiserum (1:1000), obtained as described in example I ("screening of the λ gtll M. tuberculosis recombinant DNA library with anti-32-kDa antiserum") in the presence of E. coli lysate or with monoclonal

anti-hTNF-antibody which crossreacts with mTNF (Innogenetics, n° 17F5D10) for at least 2 hours on a rotating platform. After washing twice with NT buffer + 0.02% Triton.X.100, blots are incubated for at least 1 hour with the secondary antiserum phosphatase-conjugated swine anti-rabbit immunoglobulins (1/500; Prosan) in the first case, and phosphatase conjugated rabbit alkaline anti-mouse immunoglobulins (1/500; Sigma) in the second case.

Blots are washed again twice with NT buffer + 0.02% Triton X100 and visualisation is then performed with nitro blue tetrazolium (NBT) and 5-bromo-4-chloro-3-indolyl-phosphate (BCIP) from Promega using conditions recommended by the supplier.

B. RESULTS

Upon induction of K12AH cells containing pmTNF-MPH-Mt32, a clearly visible band of about 35-kDa appears on CBB stained gels, already one hour after start of induction (Fig. 14a). This band, corresponding to roughly 25% of total protein contents of the cell, reacts strongly with anti-32-kDa and anti-mTNF antisera (Fig. immunoblots 14b). However, this represents a cleavage product of the original fusion protein, since a minor band, around 37 kDa, is also visible on immunoblots, reacting specifically with both antisera as well. This suggests that extensive cleavage of the recombinant mTNF-His6-P32 takes place about 2-3 kDa from its carboxyterminal end.

EXAMPLE VII : PURIFICATION OF RECOMBINANT ANTIGEN ON IMMOBILIZED METAL ION AFFINITY CHROMATOGRAPHY (IMAC) :

The hybrid protein $mTNF-His_6-P_{32}$ (amino acid sequence, see fig. 13) expressed by $K12\Delta H$ cells containing pmTNF.MPH.Mt32, is especially designed to facilitate purification by IMAC, since the 6 successive histidines in the polylinker sequence bring about a strong affinity for metal ions (HOCHULI et al, 1988).

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a. Preparation of the crude cell extract:

12 l of E. coli cells K12 Δ H containing plasmid pmTNF-MPH-Mt32 were grown in Luria Broth containing tetracycline (10 μ g/ml) at 28°C to an optical density (600 nm) of 0.2 and then induced by shifting the temperature to 42°C. After 3 hours of induction, cells were harvested by centrifugation (Beckman, JA 10 rotor, 7.500 rpm, 10 min). The cell paste was resuspended in lysis buffer (10 mM KCl, 10 mM Tris-HCl pH 6.8, 5 mM EDTA) to a final concentration of 50% (w/v) cells.

 ϵ -NH₂-capronic acid and dithiotreitol (DTT) were added to a final concentration of resp. 20 mM and 1 mM, to prevent proteolytic degradation. This concentrated cell suspension was stored overnight at -70°C.

Cells were lysed by passing them three times through a French press (SLM-Aminco) at a working pressure of 800-1000 psi. During and after lysis, cells were kept systematically on ice.

The cell lysate was cleared by centrifugation (Beckman, JA 20, 18.000 rpm, 20 min, 4°C). The supernatant (SN) was carefully taken off and the pellet, containing membranes and inclusion bodies, was kept for further work since preliminary experiments had shown that the protein was mainly localised in the membrane fraction.

7 M guanidinium hydrochloride (GuHCl, marketed by ICN) in 100 mM phosphate buffer pH 7.2 was added to the pellet volume to a final concentration of 6 M GuHCl. The pellet was resuspended and extracted in a bounce tissue homogenizer (10 cycles).

After clearing (Beckman, JA 20, 18.000 rpm, 20 min, 4°C), about 100 ml of supernatant was collected (= extract 1) and the removing pellet was extracted again as described above (= extract 2, 40 ml).

The different fractions (SN,EX1,EX2) were analysed on SDS-PAGE (Laemmli, Nature 1970; 227:680) together

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with control samples of the induced culture. Scanning of the gel revealed that the recombinant protein makes up roughly 25% of the total protein content of the induced cell culture. After fractionation most of the protein was found back in the extracts. No difference was noticed between reducing and non-reducing conditions (plus and minus β -mercaptoethanol).

b. <u>Preparation of the Ni⁺⁺ IDA (Imino diacetic</u> acid) column:

5 ml of the chelating gel, Chelating Sepharose 6B (Pharmacia) is washed extensively with water to remove the ethanol in which it is stored and then packed in a "Econo-column" (1 x 10 cm, Biorad). The top of the column is connected with the incoming fluid (sample, buffer, etc) while the end goes to the UV₂₈₀ detector via a peristaltic jump. Fractions are collected using a fraction collector and, when appropriate, pH of the fractions is measured manually.

The column is loaded with Ni⁺⁺ (6 ml NiCl₂.6H₂O; 5 μ g/ μ l) and equilibrated with starting buffer (6 M guanidinium hydrochloride, 100 mM phosphate buffer, pH 7.2).

After having applied the sample, the column is washed extensively with starting buffer to remove unbound material.

To elute the bound material, 2 different elution procedures are feasible:

- 1) elution by decreasing pH,
- 2) elution by increasing imidazol concentration. Both will be discussed here.

To regenerate the column, which has to be done after every 2-3 runs, 20 ml (about 5 column volumes) of the following solutions are pumped successively through the column:

- 0.05 M EDTA 0.5 M NaCl
- 0.1 M NaOH

- H₂O

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- 6 ml $NiCl_2.6H_2O$ (5 mg/ml).

After equilibrating with starting buffer the column is ready to use again.

c. Chromatography:

M guanidinium contained 6 buffers All hydrochloride throughout the chromatography. The column was developed at a flow rate of 0.5 ml/min at ambient temperature. Fractions of 2 ml were collected and, when SDS-PAGE by analysed further appropriate, Coomassie with were stained Gels immunoblotting. Brilliant Blue R250 and silver stain, as described by ANSORGE (1985). Immunoblotting was carried out as described in example I. The primary antiserum used was anti-32kDa-antiserum polyclonal either obtained as described in example I ("screening of the Agt11 M. tuberculosis recombinant DNA library with anti-32kDa-antiserum") or anti-E. coli-immunoglobulins (1/500; PROSAN), or monoclonal anti-hTNF-antibody which cross-reacts with mTNF (Innogenetics, N° 17F5D10). The secondary antiserum was alkaline phosphatase conjugated swine anti-rabbit immunoglobulins (1/500, PROSAN), or phosphatase conjugated rabbit-anti-mouse alkaline immunoglobulins (1/500, Sigma).

C1. Elution with decreasing pH :

Solutions used:

A: 6 M GuHCl 100 mM phosphate pH 7.2

B: 6 M GuHCl 25 mM phosphate pH 7.2

C: 6 M GuHCl 50 mM phosphate pH 4.2

After applying 3 ml of extract 1 ($OD_{280} = 32.0$) and extensively washing with solution A, the column is equilibrated with solution B and then developed with a linear pH gradient from 7.2 to 4.2 (25 ml of solution B and 25 ml of solution C were mixed in a gradient former). The elution profile is shown in figure 15.

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From SDS-PAGE analysis (Coomassie and silverstain) it was clear that most of the originally bound recombinant protein was eluted in the fractions between pH 5.3 and 4.7.

Screening of these fractions on immunoblot with anti-32-kDa and the 17F5D10 monoclonal antibody showed that, together with the intact recombinant protein, also some degradation products and higher aggregation forms of the protein were present, although in much lower amount. Blotting with anti-E. coli antibody revealed that these fractions (pH 5.3-4.7) contained immunodetectable contaminating E. coli proteins (75, 65, 43, 35 and 31 kDa bands) lipopolysaccharides..

C2. Elution with increasing imidazol concentration:

Solutions used :

A: 6 M GuHCl 100 mM phosphate pH 7.2

B: 6 M GuHCl 50 mM imidazol pH 7.2

C: 6 M GuHCl 100 mM imidazol pH 7.2

D: 6 M GuHCl 15 mM imidazol pH 7.2

E: 6 M GuHCl 25 mM imidazol pH 7.2

F: 6 M GuHCl 35 mM imidazol pH 7.2

Sample application and washing was carried out as in C1, except that after washing, no equilibration was necessary with 6 M GuHCl 25 mM phosphate. The column was first developed with a linear gradient of imidazol going from 0 to 50 mM (25 ml of solution A and 25 ml of solution B were mixed in a gradient former) followed by a step elution to 100 mM imidazol (solution C). During the linear gradient, proteins were gradually eluted in a broad smear, while the step to 100 mM gave rise to a clear peak (fig. 16).

SDS-PAGE analysis of the fractions revealed that in the first part of the linear gradient (fr 1-24) most

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contaminating <u>E. coli</u> proteins were washed out, while the latter part of the gradient (fr 25-50) and the 100 mM peak contained more than 90% of the recombinant protein.

As in C1, these fractions showed, besides a major band of intact recombinant protein, some minor bands of degradation and aggregation products. However, in this case, the region below 24-kDa seemed nearly devoid of protein bands, which suggests that less degradation products co-elute with the intact protein. Also, the same contaminating <u>E. coli</u> proteins were detected by immunoblotting, as in C1, although the 31-kDa band seems less intense and even absent in some fractions.

In a second stage, we developed the column with a step gradient of increasing imidazol concentrations. After having applied the sample and washed the column, 2 column volumes (about 8 ml) of the following solutions were brought successively onto the column: solution D, E, F and finally 4 column volumes of solution C. The stepgradient resulted in a more concentrated elution profile (fig. 17) which makes it more suitable for scaling up purposes.

In conclusion, the mTNF-His $_6$ -P $_{32}$ protein has been purified to at least 90% by IMAC. Further purification can be achieved through a combination of the following purification steps:

- IMAC on chelating superose (Pharmacia)
- ion exchange chromatography (anion or cation)
- reversed phase chromatography
- gel filtration chromatography
- immunoaffinity chromatography
- elution from polyacrylamide gel.

These chromatographic methods are commonly used for protein purification.

The plasmids of figures 10b, 11b and 12b are new.

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CLAIMS

- 1. Recombinant polypeptide containing in its polypeptidic chain, one at least of the following amino acid sequences:
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 3a and fig. 3b, or
- the one extending from the extremity constituted by amino acid at position (275) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,

and the peptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids in so far as this modification does not alter the following properties:

the polypeptides react with rabbit polyclonal antiserum raised against the protein of 32-kDa of $\underline{\text{M.}}$ bovis BCG culture filtrate, and/or

react selectively with human sera from tuberculosis patients and particularly patients developing an evolutive tuberculosis at an early stage,

and/or react with the amino acid sequence extending from the extremity constituted by amino acid at position (1), to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b.

- 2. Recombinant polypeptide according to claim 1, containing in its polypeptidic chain, one at least of the following amino acid sequences:
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 4a and fig. 4b, or

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- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 4a and fig. 4b, or

- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 4a and fig. 4b, or
- the one extending from the extremity constituted by amino acid at position (275) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,

and the peptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids in so far as this modification does not alter the following properties:

the polypeptides react with rabbit polyclonal antiserum raised against the protein of 32-kDa of $\underline{\text{M.}}$ bovis BCG culture filtrate, and/or

react selectively with human sera from tuberculosis patients and particularly patients developing an evolutive tuberculosis at an early stage,

and/or react with the amino acid sequence extending from the extremity constituted by amino acid at position (1), to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b.

3. Recombinant polypeptide according to claim 1, containing in its polypeptidic chain, one at least of the following amino acid sequences:

- the one extending from the extremity constituted by amino acid at position (-30) to the extremity constituted by amino acid at position (-1) represented on fig. 5, or

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- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 5, or
- the one extending from the extremity constituted by amino acid at position (275) to the extremity constituted by amino acid at position (295) represented on fig. 5,
- and the peptidic sequences resulting from the modification by substitution and/or by addition and/or by deletion of one or several amino acids in so far as

this modification does not alter the following properties:

the polypeptides react with rabbit polyclonal antiserum raised against the protein of 32-kDa of <u>M. bovis</u> BCG culture filtrate, and/or

react selectively with human sera from tuberculosis patients and particularly patients developing an evolutive tuberculosis at an early stage,

and/or react with the amino acid sequence extending from the extremity constituted by amino acid at position (1), to the extremity constituted by amino acid at position (295) represented on fig. 5.

- 4. Recombinant polypeptide according to claim 1, containing in its polypeptidic chain, one at least of the following amino acid sequences:
- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-49) to to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,

- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,

- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,

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- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b.
- 5. Recombinant polypeptide according to claim 2, containing in its polypeptidic chain, one at least of the following amino acid sequences:
- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,

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- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-49) to to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,

- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b.
- 6. Recombinant polypeptide according to claim 3, containing in its polypeptidic chain, one at least of the following amino acid sequences:
- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (295) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-43) to the extremity constituted by amino acid at position (-1) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-30) to the extremity constituted by amino acid at position (295) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-43) to the extremity constituted by amino acid at position (295) represented on fig. 5.
- 7. Recombinant polypeptide according to claim 1, consisting in one of the following amino acid sequences:
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity

constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,

- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity

constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,

- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (-1) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (294) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 3a and fig. 3b,
- the one extending from the extremity constituted by amino acid at position (275) to the extremity

constituted by amino acid at position (294) represented on fig. 3a and fig. 3b.

- 8. Recombinant polypeptide according to claim 2, consisting in one of the following amino acid sequences:
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-59) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-55) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-49) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-47) to the extremity

constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,

- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-42) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (-29) to the extremity constituted by amino acid at position (-1) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (101) to the extremity

constituted by amino acid at position (120) represented on fig. 4a and fig. 4b,

- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 4a and fig. 4b,
- the one extending from the extremity constituted by amino acid at position (275) to the extremity constituted by amino acid at position (294) represented on fig. 4a and fig. 4b.
- 9. Recombinant polypeptide according to claim 3, consisting in one of the following amino acid sequences:
- the one extending from the extremity constituted by amino acid at position (-43) to the extremity constituted by amino acid at position (295) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-43) to the extremity constituted by amino acid at position (-1) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-30) to the extremity constituted by amino acid at position (295) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (-30) to the extremity constituted by amino acid at position (-1) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (1) to the extremity constituted by amino acid at position (295) represented on fig. 5,

- the one extending from the extremity constituted by amino acid at position (12) to the extremity constituted by amino acid at position (31) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (36) to the extremity constituted by amino acid at position (55) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (77) to the extremity constituted by amino acid at position (96) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (101) to the extremity constituted by amino acid at position (120) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (175) to the extremity constituted by amino acid at position (194) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (211) to the extremity constituted by amino acid at position (230) represented on fig. 5,
- the one extending from the extremity constituted by amino acid at position (275) to the extremity constituted by amino acid at position (295) represented on fig. 5.
- 10. Amino acid sequences constituted by a polypeptide according to claims 1 to 9, and a protein or an heterologous sequence with respect to said polypeptide, said protein or heterologous sequence comprising from about 1 to about 1000 amino acids.
- 11. Amino acid sequence according to claim 10, wherein the heterologous protein is β -galactosidase.
 - 12. Nucleic acid comprising

- a nucleotide sequence coding for anyone of the polypeptides according to claims 1 to 11,
- or nucleotide sequences which hybridize with the nucleotide sequences coding for anyone of the polypeptides according to claims 1 to 11,
- or nucleotide sequences which are complementary to the nucleotide sequences coding for any of the polypeptides according to claims 1 to 11,
- the above mentioned nucleotide sequences wherein T can be replaced by U.
- 13. Nucleic acid according to claim 12, comprising one at least of the following nucleotide sequences:
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (182) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1242) to the extremity constituted by nucleotide at position (1358), wherein N represents one of the five A, T, C, G or I nucleotides, represented in fig. 3a and fig. 3b,
- or above said nucleotide sequences wherein T is replaced by U,
- or nucleic acids which hybridize with said above mentioned nucleotide sequences or the complementary sequences thereof.
- 14. Nucleic acid according to claim 13, comprising one at least of the following nucleotide sequences:

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (182) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1242) to the extremity constituted by nucleotide at position (1358), wherein N represents one of the five A, T, C, G or I nucleotides, represented in fig. 4a and fig. 4b,
- or above said nucleotide sequences wherein T is replaced by U,
- or nucleic acids which hybridize with said above mentioned nucleotide sequences or the complementary sequences thereof.
- 15. Nucleic acid according to claim 13, comprising one at least of the following nucleotide sequences:
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (219) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (220) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1104) to the extremity constituted by nucleotide at position (1299),

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or above said nucleotide sequences wherein T is replaced by U,

or nucleic acids which hybridize with said above mentioned nucleotide sequences or the complementary sequences thereof.

- 16. Nucleic acid according to claim 13, comprising one of the following sequences:
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (194) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (212) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (218) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (272) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b.

- 17. Nucleic acid according to claim 14, comprising one of the following sequences:
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (194) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (212) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (218) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (272) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity

constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity

constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,

- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b.
- 18. Nucleic acid according to claim 15, comprising one of the following sequences:
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (129) represented in fig. 5,

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- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (219) represented in fig. 5, - the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1104) represented in fig. 5, - the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1299) represented in fig. 5, - the one extending from the extremity constituted by the (90) to nucleotide position at constituted by nucleotide at position (219) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (90) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (90) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (220) to the extremity constituted by nucleotide at position (1299) represented in fig. 5.
- 19. Nucleic acid according to claim 13, consisting in one of the following nucleotide sequences:

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- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (182) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (194) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (212) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (218) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (272) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,

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- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,

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- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (359) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1241) represented in fig. 3a and fig. 3b,

- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b,
- the one extending from the extremity constituted by nucleotide at position (1242) to the extremity constituted by nucleotide at position (1358) represented in fig. 3a and fig. 3b.
- 20. Nucleic acid according to claim 14, consisting in one of the following nucleotide sequences:
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (182) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (194) represented in fig. 4a and fig. 4b.
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (212) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (218) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (272) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted

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by nucleotide at position (1241) represented in fig. 4a and fig. 4b,

- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (183) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (195) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,

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- the one extending from the extremity constituted by nucleotide at position (213) to the extremity

constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,

- the one extending from the extremity constituted by nucleotide at position (213) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (219) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (234) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (359) represented in fig. 4a and fig. 4b,

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- the one extending from the extremity constituted by nucleotide at position (273) to the extremity

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constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,

- the one extending from the extremity constituted by nucleotide at position (273) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1241) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (360) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b,
- the one extending from the extremity constituted by nucleotide at position (1242) to the extremity constituted by nucleotide at position (1358) represented in fig. 4a and fig. 4b.
- 21. Nucleic acid according to claim 15, consisting in one of the following nucleotide sequences:
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (129) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (219) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,

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- the one extending from the extremity constituted by nucleotide at position (90) to the extremity constituted by nucleotide at position (219) represented in fig. 5,

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- the one extending from the extremity constituted by nucleotide at position (90) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (90) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (219) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (130) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (220) to the extremity constituted by nucleotide at position (1104) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (220) to the extremity constituted by nucleotide at position (1299) represented in fig. 5,
- the one extending from the extremity constituted by nucleotide at position (1104) to the extremity constituted by nucleotide at position (1299) represented in fig. 5.
- 22. Recombinant nucleic acid containing at least one of the nucleotide sequences according to claims 13 to 21, inserted in a heterologous nucleic acid.

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23. DNA or RNA primer constituted by one of the following sequences:

- A(i) CAGCTTGTTGACAGGGTTCGTGGC
- A(ii) GGTTCGTGGCGCCGTCACG
- A(iii) CGTCGCGCGCCTAGTGTCGG
- A(iv) CGGCGCCGGTCGGTGGCACGGCGA
- A(v) CGTCGGCGCGCCCTAGTGTCGG
- B TCGCCCGCCCTGTACCTG
- C GCGCTGACGCTGGCGATCTATC
- D CCGCTGTTGAACGTCGGGAAG
- E AAGCCGTCGGATCTGGGTGGCAAC
- F(i) ACGGCACTGGGTGCCACGCCCAAC
- F(ii) ACGCCCAACACCGGGCCCGCCA
- F(iii) ACGGGCACTGGGTGCCACGCCCAAC
- F(iv) ACGCCCAACACCGGGCCCGCGCCCCA
- 24. DNA or RNA primer set constituted by any of the nucleotide sequences A(i), A(ii), A(iii), A(iv), A(v), A(v),
- A(i), A(ii), A(iii), A(iv), A(v), B, C, D, E, F(i), F(ii), F(iii) and F(iv) having the meaning of claim 11, and

advantageously constituted by the following elements:

- A(i)
- or A(ii)
- or A(iii) and the complement of B
- or A(iv)
- or A(v)

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A(i)
or A(ii)
                 and the complement of C
or A(iii)
or A(iv)
or A(v)
                  and the complement of C
  В
  A(i)
or A(ii)
                and the complement of F
or A(iii)
or A(iv)
or A(v)
  A(i)
or A(ii)
                and the complement of D
or A(iii)
or A(iv)
or A(v)
   A(i)
or A(ii)
               and the complement of E
or A(iii)
or A(iv)
or A(v)
                 and the complement of D
   В
                  and the complement of E
   В
                  and the complement of F
   В
                  and the complement of D
   C
                  and the complement of E
   C
                  and the complement of F
   С
                  and the complement of E
   D
                  and the complement of F
   D
                  and the complement of F.
   E
     25. Recombinant vector, particularly for cloning
        expression, comprising a vector
                                               sequence,
and/or
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notably of the type plasmid, cosmid or phage, and a recombinant nucleic acid according to anyone of claims 13 to 21, in one of the non essential sites for its replication.

- 26. Recombinant vector according to claim 25, containing in one of its non essential sites for its replication necessary elements to promote the expression of polypeptides according to anyone of claims 1 to 12 in a cellular host and possibly a promoter recognized by the polymerase of the cellular host, particularly an inductible promoter and possibly a signal sequence and/or an anchoring sequence.
- 27. Recombinant vector according to claim 26, containing the elements enabling the expression by \underline{E} . \underline{coli} of a nucleic acid according to anyone of claims 6 to 9 inserted in the vector, and particularly the elements enabling the expression of the gene or part thereof of β -galactosidase.
- 28. Cellular host which is transformed by a recombinant vector according to anyone of claims 25 to 27, and comprising the regulation elements enabling the expression of the nucleotide sequence coding for the polypeptide according to anyone of claims 1 to 12 in this host.
- 29. Cellular host according to claim 28, chosen from among bacteria such as \underline{E} . \underline{coli} , transformed by the vector according to claim 25, or chosen from among eukaryotic organism, transformed by the vector according to claim 25.
- 30. Expression product of a nucleic acid expressed by a transformed cellular host according to anyone of claims 28 or 29.
- 31. Antibody characterized by the fact that it is directed against a recombinant polypeptide according to anyone of claims 1 to 12.

32. Nucleotidic probes, hybridizing with anyone of the nucleic acids according to claims 13 to 21 or with their complementary sequences, and particularly the probes chosen among the following nucleotidic sequences

Probes A(i), A(ii), A(iii) and A(iv)

- A(i) CAGCTTGTTGACAGGGTTCGTGGC
- A(ii) GGTTCGTGGCGCCGTCACG
- A(iii) CGTCGCGCGCCTAGTGTCGG
- A(iv) CGGCGCCGTCGGTGGCACGGCGA
- A(v) CGTCGGCGCGCCCTAGTGTCGG

Probe B

TCGCCCGCCCTGTACCTG

Probe C

GCGCTGACGCTGGCGATCTATC

Probe D

CCGCTGTTGAACGTCGGGAAG

Probe E

AAGCCGTCGGATCTGGGTGGCAAC

Probes F(i) and F(ii)

- F(i) ACGGCACTGGGTGCCACGCCCAAC
- F(ii) ACGCCCAACACCGGGCCCGCCA
- F(iii) ACGGGCACTGGGTGCCACGCCCAAC
- F(iv) ACGCCCCAACACCGGGCCCGCGCCCCA
- or their complementary nucleotidic sequences.
- 33. Process for preparing a recombinant polypeptide according to anyone of claims 1 to 12 comprising the following steps:

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- the culture in an appropriate medium of a cellular host which has previously been transformed by an appropriate vector containing a nucleic acid according to anyone of claims 12 to 22, and
- the recovery of the polypeptide produced by the above said transformed cellular host from the above said culture medium.
- 34. Method for the <u>in vitro</u> diagnostic of tuberculosis in a patient liable to be infected by Mycobacterium tuberculosis comprising the following steps:
- the possible previous amplification of the amount of the nucleotide sequences according to anyone of claims 12 to 22, liable to be contained in a biological sample taken from said patient by means of a DNA primer set according to claim 24,
- contacting the above mentioned biological sample with a nucleotide probe according to claim 32, under conditions enabling the production of an hybridization complex formed between said probe and said nucleotide sequence,
- detecting the above said hybridization complex which has been possibly formed.
- 35. Method for the <u>in vitro</u> diagnostic of tuberculosis in a patient liable to be infected by Mycobacterium tuberculosis comprising
- contacting a biological sample taken from a patient with a polypeptide according to anyone of claims 1 to 11, under conditions enabling an <u>in vitro</u> immunological reaction between said polypeptide and the antibodies which are possibly present in the biological sample and the <u>in vitro</u> detection of the antigen/antibody complex which has been possibly formed.
- 36. Method for the <u>in vitro</u> diagnostic of tuberculosis in a patient liable to be infected by <u>M.</u> tuberculosis, comprising the following steps:

- contacting the biological sample with an appropriate antibody according to claim 31, under conditions enabling an <u>in vitro</u> immunological reaction between said antibody and the antigens of <u>M. tuberculosis</u> which are possibly present in the biological sample and the <u>in vitro</u> detection of the antigen/antibody complex which may be formed.
- 37. Necessary or kit for an <u>in vitro</u> diagnostic method of tuberculosis in a patient liable to be infected by Mycobacterium tuberculosis according to claim 34, comprising
- a determined amount of a nucleotide probe according to claim 32,
- advantageously the appropriate medium for creating an hybridization reaction between the sequence to be detected and the above mentioned probe,
- advantageously, reagents enabling the detection of the hybridization complex which has been formed between the nucleotide sequence and the probe during the hybridization reaction.
- 38. Necessary or kit for an <u>in vitro</u> diagnostic method of tuberculosis in a patient liable to be infected by Mycobacterium tuberculosis according to claim 35, comprising
- a polypeptide according to anyone of claims 1 to 11,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complex which has been produced by the immunological reaction, said reagents possibly having a label, or being liable to be recognized by a labeled reagent, more particularly in the case where the above mentioned polypeptide is not labeled.
- 39. Necessary or kit for an <u>in vitro</u> diagnostic method of tuberculosis in a patient liable to be

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infected by Mycobacterium tuberculosis according to claim 36, comprising

- an antibody according to claim 31,
- reagents for making a medium appropriate for the immunological reaction to occur,
- reagents enabling to detect the antigen/antibody complexes which have been produced by the immunological reaction, said reagents possibly having a label or being liable to be recognized by a label reagent, more particularly in the case where the above mentioned antibody is not labeled.
- 40. Immunogenic composition comprising a polypeptide according to anyone of claims 1 to 11, in association with a pharmaceutically acceptable vehicle.
- 41. Vaccine composition comprising among other immunogenic principles anyone of the polypeptides according to claims 1 to 11 or the expression product of claim 30, possibly coupled to a natural protein or to a synthetic polypeptide having a sufficient molecular weight so that the conjugate is able to induce in vivo the production of antibodies neutralizing Mycobacterium tuberculosis, or induce in vivo a cellular immune response by activating M. tuberculosis antigen-responsive T cells.
- 42. Process for the enzymatical amplification of a nucleotide sequence according to claims 12 to 22, and detection of the amplified nucleotide sequence, wherein the amplification is achieved by PCR technique by means of a primer set and the detection of the PCR amplified product is achieved by a hybridization reaction with a detection probe constituted by an oligonucleotide sequence of at least 10 nucleotides, said sequence being located between the two PCR primers which have been used for amplifying said nucleotide sequence,

- the primer set and detection probe used being preferably chosen among the following elements:

Primer set

P1 GAGTACCTGCAGGTGCCGTCGCTCGATGGGCCG

P2 compl. GTACCACTCGAACGCCGGGGTGTTGAT

Probe B

TCGCCCGCCCTGTACCTG

Primer set

P1 GAGTACCTGCAGGTGCCGTCGCCGTCGATGGGCCG

P3 compl. TCCCACTTGTAAGTCTGGCA

Probe B

TCGCCCGCCCTGTACCTG

Primer set

P1 GAGTACCTGCAGGTGCCGTCGCTCGATGGGCCG

P4 compl. CGGCAGCTCGCTGGTCAGGA

Probe B

TCGCCCGCCCTGTACCTG

Primer set

P1 GAGTACCTGCAGGTGCCGTCGCTCGATGGGCCG

P5 compl. GCGTCACCCATCGCCAGGCCGATCAGG

Probe B

TCGCCCGCCCTGTACCTG or

Probe C

GCGCTGACGCTGGCGATCTATC

Primer set

P1 GAGTACCTGCAGGTGCCGTCGCCGTCGATGGGCCG

P6 compl. GCGCCCCAGTACTCCCAGCTGTGCGT

Probe B

TCGCCCGCCCTGTACCTG or

Probe C

GCGCTGACGCTGGCGATCTATC or

Probe D

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CCGCTGTTGAACGTCGGGAAG or

Probe E

AAGCCGTCGGATCTGGGTGGCAAC

Primer set

P2 ATCAACACCCCGGCGTTCGAGTGGTAC

P5 compl. GCGTCACCCATCGCCAGGCCGATCAGG

Probe C

GCGCTGACGCTGGCGATCTATC

Primer set

P2 ATCAACACCCCGGCGTTCGAGTGGTAC

P6 compl. GCGCCCCAGTACTCCCAGCTGTGCGT

Probe C

GCGCTGACGCTGGCGATCTATC or

Probe D

CCGCTGTTGAACGTCGGGAAG or

Probe E

AAGCCGTCGGATCTGGGTGGCAAC

Primer set

P3 TGCCAGACTTACAAGTGGGA

P5 compl. GCGTCACCCATCGCCAGGCCGATCAGG

Probe C

GCGCTGACGCTGGCGATCTATC

Primer set

P3 TGCCAGACTTACAAGTGGGA

P6 compl. GCGCCCCAGTACTCCCAGCTGTGCGT

Probe C

GCGCTGACGCTGGCGATCTATC or

Probe D

CCGCTGTTGAACGTCGGGAAG or

Probe E

AAGCCGTCGGATCTGGGTGGCAAC

Primer set

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P4 TCCTGACCAGCGAGCTGCCG

P5 compl. GCGTCACCCATCGCCAGGCCGATCAGG

Probe C

GCGCTGACGCTGGCGATCTATC

Primer set

P4 TCCTGACCAGCGAGCTGCCG

P6 compl. GCGCCCCAGTACTCCCAGCTGTGCGT

Probe C

GCGCTGACGCTGGCGATCTATC or

Probe D

CCGCTGTTGAACGTCGGGAAG or

Probe E

AAGCCGTCGGATCTGGGTGGCAAC

Primer set

P5 CCTGATCGGCCTGGCGATGGGTGACGC

P6 compl. GCGCCCCAGTACTCCCAGCTGTGCGT

Probe D

CCGCTGTTGAACGTCGGGAAG or

Probe E

AAGCCGTCGGATCTGGGTGGCAAC

or the primer set being preferably chosen among the primer sets according to claim 24, and the detection probe being constituted by any oligonucleotide sequence of at least 10 nucleotides, said sequence being located between the two PCR primers constituting the primer set which has been used for amplifying said nucleotide sequence.

43. A vector sequence forming part of a recombinant vector according to claim 25, said vector sequence having either the nucleic acid sequence represented in fig. 10b, or the nucleic acid sequence represented in fig. 11b, or the nucleic acid sequence represented in fig. 12b.

- 44. Plasmids comprising either the nucleic acid sequence of fig. 10b, or the nucleic acid sequence of fig. 11b, or the nucleic acid sequence of fig. 12b.
- 45. Peptides of claim 1, advantageously used to produce antibodies, particularly monoclonal antibodies and which have the following amino acid sequences:

Amino acid		Amino acid		
position		position		
(NH ₂ -terminal	(COOH-terminal)			
12	OUDCDGWGDDTIGIOGOGO			
12	QVPSPSMGRDIKVQFQSGGA	31		
36	LYLLDGLRAQDDFSGWDINT	55		
77	SFYSDWYQPACRKAGCQTYK	96		
101	LTSELPGWLQANRHVKPTGS	120		
175	KASDMWGPKEDPAWQRNDPL	194		
211	CGNGKPSDLGGNNLPAKFLE	230		
275	KPDLQRHWVPRPTPGPPQGA	294		
77	SFYSDWYQPACGKAGCQTYK	96		
276	PDLORALGATPNTGPAPOGA	299		

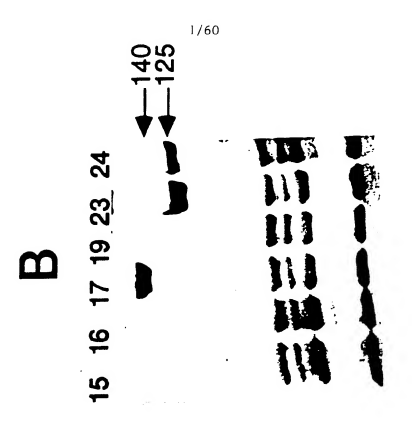
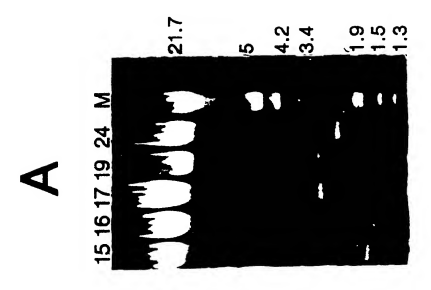
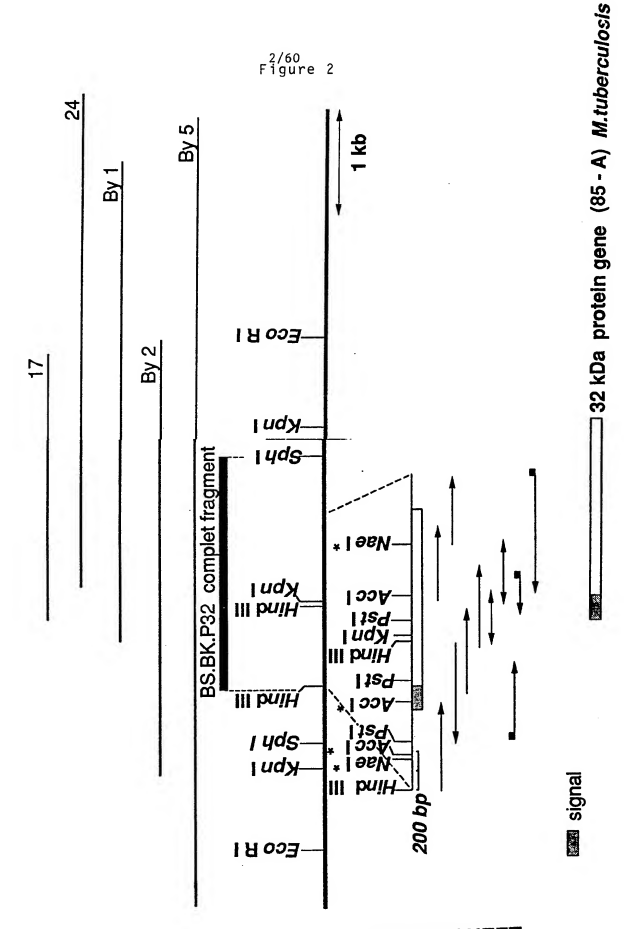


Figure |



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CGACACATGCCCAGACACTGCGGAAATGCCACCTTCAGGCCGTCGCGTCGGT cccgaa *| trggc |*cgtgaacgaccgccgg | a taa | gggtttcggcggtgcgcttgatgcgggt

GGACGCCC AGGTTGTGGTTGACTACACGAGCACTGCCGGGCCCCAGCGCCTGCAGTCTGACCT AATTCAGGATGCGCCCAAACATGCATGCG TTGAGA TGAGATTGAGGATGAGA MET-ARG-PRO-ASN-MET-HIS- GLY-CYS-VAL- GLU- MET- ARG-MET-ARG-GLU-ALA-ARG

<u>a r|g-cag-ctt-gtt-gac-agg-gtt-cgt-ggc-gcc-gtc-acg-ggt-alg-tcg-cgt-cga-ctc-gtg-gtc-</u> -47 -49

MET-GLN-LEU-VAL-ASP-ARG-VAL-ARG-GLY-ALA-VAL-THR-GLY-MET-SER-ARG-ARG-LEU-VAL-VAL-GGG-GCC-GIC-XCG-CXC-XAA-GIG-ICG-GGI-CIG-GIC-GCC-GCC-GIC-GGI-GGC-ACG-GCG-ACG-ACC-GCG--42

3/60 ${\tt GLY-ALA-VAL-a_1-b_1-LEU-VAL-SER-GLY-LEU-VAL-GLY-ALA-VAL-GLY-GLY-THR-ALA-THR-ALA-GLY-ALA-VAL-GLY-GLY-THR-ALA-THR-A$

GLY-ALA-phe-ser-arg-pro-gly-leu-pro-val-glu-tyr-leu-gln-val-pro-ser-pro-ser-met-GGG-GCA-TTT-TCC-CGG-CCG-GGC-TTG-CCG-GTG-GAG-TAC-CTG-CAG-GTG-CCG-TCG-CCG-TCG-ATG-354

GGC-CGT-GAC-ATC-AAG}-GTC-CAA-TTC-CAA-AGT-GGT-GGT-GCC-AAC-TCG-CCC-GCC-CTG-TAC-CTGgly-arg-asp-ile-lys -val-gln-phe-gln-ser-gly-gly-ala-asn-ser-pro-ala-leu-tyr-leu-

† 17

CTC-GAC-GGC-CTG+CGC-GCG-CAG-GAC-GAC-TTC-AGC-GGC-TGG-GAC-ATC-AAC-ACC-CCG-GCG-TTCleu-asp-gly-leu-arg-ala-gln-asp-asp-phe-ser-gly-trp-asp-ile-asn-thr-pro-ala-phe-

GAG-TGG-TAC-GAC-CAG-TCG-GGC-CTG-TCG-GTG-GTC-ATG-CCG-GTG-GGT-GGC-CAG-TCA-AGC-TTCglu-trp-tyr-asp-gln-ser-gly-leu-ser-val-val-met-pro-val-gly-gly-gln-ser-ser-phe-534

tyr-ser-asp-trp-tyr-gln-pro-ala-cys- a, -lys-ala-gly-cys-gln- thr-tyr-lys-trp-glu-TAC-TCC-GAC-TGG-TAC-CAG-CCC-GCC-TGC-ZGC-AAG-GCC-GGT-TGC-CAG-(ACT-TAC-AAG-TGG-GAG-

Figure

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- ACC-TTC-CTG-ACC-AGC-GAG-CTG-CCG-GGG-TGG-CTG-CAG-GCC-AAC-AGG-CAC-GTC-AAG-CCC-ACCthr-phe-leu-thr-ser-glu-leu-pro-gly-trp-leu-gln-ala-asn-arg-his-val-lys-pro-thr-
- gly-ser-ala-val-val-gly-leu-ser-met-ala-ala-ser-ser-ala-leu-thr-leu-ala-ile-tyr-GGA-AGC-GCC-GTC-GTC-GGT-CTT-TCG-ATG-GCT-GCT-TCT-TCG-GCG-CTG-ACG-CTG-GCG-ATC-TAT-
 - CAC-CCC-CAG-CAG-TTC-GTC-TAC-GCG-GGA-GCG-ATG-TCG-GGC-CTG-TTG-GAC-CCC-TCC-CAG-GCG-
- his-pro-gln-gln-phe-val-tyr-ala-gly-ala-met-ser-gly-leu-leu-asp-pro-ser-gln-ala-ATG-GGT-CCC-ACC-CTG-ATC-GGC-CTG-GCG-ATG-GGT-GAC-GCT-GGC-GGC-TAC-AAG-GCC-TCC-GAC-139
 - met-gly-pro-thr-leu-ile-gly-leu-ala-met-gly-asp-ala-gly-gly-tyr-lys-ala-ser-asp-834 159
- ATG-TGG-GGC-CCG-AAG-GAG-GAC-CCG-GCG-TGG-CAG-CGC-AAC-GAC-CCG-CTG-TTG-AAC-GTC-GGG-894
- met-trp-gly-pro-lys-glu-asp-pro-ala-trp-gln-arg-asn-asp-pro-leu-leu-asn-val-gly-& 179
- lys-leu-ile-ala-asn-asn-thr-arg-val-trp-val-tyr-cys-gly-asn-gly-lys-pro-ser-asp-AAG-CTG-ATC-GCC-AAC-AAC-ACC-CGC-GTC-TGG-GTG-TAC-TGC-GGC-AAC-GGC-AAG-CCG-TCG-GAT-199 954
- leu-gly-gly-asn-asn-leu-pro-ala-lys-phe-leu-glu-gly-phe-val-arg-thr-ser-asn-ile-CTG-GGT-GGC-AAC-AAC-CTG-CCG-GCC-AAG-TTC-CTC-GAG-GGC-TTC-GTG-CGG-ACC-AGC-AAC-ATC-1014
- lys-phe-gln-asp-ala-tyr-asn-ala-gly-gly- $^{\mathrm{a}_2}$ -his-asn-gly-val-phe-asp-phe-pro-asp-AAG-TIC-CAA-GAC-GCC-TAC-AAC-GCC-GGT- GGW-ZGC -CAC-AAC-GGC-GTG-TIC-GAC-TIC-CCG-GAC-1074
- AGC-GGT-ACG-CAC-AGC-TGG-GAG-TAC-TGG-GGC-GCG-CAG-CTC-AAC-GCT-ATG-AAG-CCC-GAC-CTGser-gly-thr-his-ser-trp-glu-tyr-trp-gly-ala-gln-leu-asn-ala-met-lys-pro-asp-leu-1134
- CAA-CG -CAC-TGG-GTG-CCA-CGC-CCA-ACA-CCG-GGC-CCG-KCL-CAG-GGC-GCC-TAGCTCCGAACAGACA $\operatorname{gln-arg-a_3} - \operatorname{b_3} - \operatorname{c_3} - \operatorname{d_3} - \operatorname{e_3} - \operatorname{f_3} - \operatorname{thr-} \operatorname{a_4} - \operatorname{gly-pro-a_5} - \operatorname{gln-gly-ala-TER}$ 1194
- CAACATCTAGCNNCGGTGACCCTTGTGGNNCANATGTTTCCTAAATCCCGTCCCTAGCTCCCGCNGCNNCCGTGTGGTTA GCTACCTGACNNCATGGGTTT 1358

CCCGAA TTGGC CGTGAACGACCGCCGG ATAA GGGTTTCGGCGGTGCGCTTGATGCGGGT CGACACATGCCCAGACACTGCGGAAATGCCACCTTCAGGCCGTCGCGTCGGT

GGACGCCCAAGTTGTGGTTGACTCACGAGCACTGCCGGGCCCAGCGCCTGCAGTCTGACCT AATTCAGGATGCGCCCAAACATGCATGCGTTGAGATTGAGATTGAGATTGAGATGAGATGAAGA MET-ARG-PRO-ASN-MET-HIS- GLY-CYS-VAL- GLU- MET- ARG-MET-ARG-GLU-ALA-ARG

<u>a t | g</u>-cag-cit-git-gac-agg-git-cgt-ggc-gcc-gic-acg-ggt-<u>aig</u>-icg-cgt-cga-cic-gig-gic-MET-GLN-LEU-VAL-ASP-ARG-VAL-ARG-GLY-ALA-VAL-THR-GLY-MET-SER-ARG-ARG-LEU-VAL-VAL--47 -49

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GLY-ALA-phe-ser-arg-pro-gly-leu-pro-val-glu-tyr-leu-gln-val-pro-ser-pro-ser-met-GGG-GCA-TII-ICC-CGG-CCG-GGC-TIG-CCG-GIG-GAG-IAC-CIG-CAG-GIG-CCG-ICG-CCG-ICG-ICG-AIG-354

GGC-CGT-GAC-ATC-AAG}-GTC-CAA-TTC-CAA-AGT-GGT-GGT-GCC-AAC-TCG-CCC-GCC-CTG-TAC-CTGgly-arg-asp-ile-lys -val-gln-phe-gln-ser-gly-gly-ala-asn-ser-pro-ala-leu-tyr-leu-414

CTC-GAC-GGC-CTG-CGC-GCG-CAG-GAC-TTC-AGC-GGC-TGG-GAC-ATC-AAC-ACC-CCG-GCG-TTC-**†** 17 474 GAG-TGG-TAC-GAC-CAG-TCG-GGC-CTG-TCG-GTG-GTC-ATG-CCG-GTG-GGT-GGC-CAG-TCA-AGC-TTCglu-trp-tyr-asp-gln-ser-gly-leu-ser-val-val-met-pro-val-gly-gly-gln-ser-ser-phe-534 59

leu-asp-gly-leu-arg-ala-gln-asp-asp-phe-ser-gly-trp-asp-ile-asn-thr-pro-ala-phe-

tyr-ser-asp-trp-tyr-gln-pro-ala-cys-arg-lys-ala-gly-cys-gln-thr-tyr-lys-trp-glu-TAC-TCC-GAC-TGG-TAC-CAG-CCC-GCC-TGC-CGC-AAG-GCC-GGT-TGC-CAG-(ACT-TAC-AAG-TGG-GAG-594

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ACC-TIC-CIG-ACC-AGC-GAG-CIG-CCG-GGG-IGG-CIG-CAG-GCC-AAC-AGG-CAC-GIC-AAG-CCC-ACCthr-phe-leu-thr-ser-glu-leu-pro-gly-trp-leu-gln-ala-asn-arg-his-val-lys-pro-thr-

gly-ser-ala-val-val-gly-leu-ser-met-ala-ala-ser-ser-ala-leu-thr-leu-ala-ile-tyr-GGA-AGC-GCC-GTC-GTC-GGT-CTT-TCG-ATG-GCT-GCT-TCT-TCG-GCG-CTG-ACG-CTG-GCG-ATC-TAT-

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his-pro-gln-gln-phe-val-tyr-ala-gly-ala-met-ser-gly-leu-leu-asp-pro-ser-gln-ala-CAC-CCC-CAG-CAG-TTC-GTC-TAC-GCG-GGA-GCG-ATG-TCG-GGC-CTG-TTG-GAC-CCC-TCC-CAG-GCG-139 774

ATG-GGT-CCC-ACC-CTG-ATC-GGC-CTG-GCG-ATG-GGT-GAC-GCT-GGC-GGC-TAC-AAG-GCC-TCC-GACmet-gly-pro-thr-leu-ile-gly-leu-ala-met-gly-asp-ala-gly-gly-tyr-lys-ala-ser-asp-834

6/60 met-trp-gly-pro-lys-glu-asp-pro-ala-trp-gln-arg-asn-asp-pro-leu-leu-asn-val-glyatg-tgg-'ggc-ccg-aag-gag-gac-ccg-gcg-tgg-cag-cgc-aac-gac-ccg-ctg-ttg-tag-gac-gtc-ggg-

AAG-CTG-ATC-GCC-AAC-ACC-CGC-GTC-TGG-GTG-TAC-TGC-GGC-AAC-GGC-AAG-CCG-TCG-GATlys-leu-ile-ala-asn-asn-thr-arg-val-trp-val-tyr-cys-gly-asn-gly-lys-pro-ser-asp-

leu-gly-gly-asn-asn-leu-pro-ala-lys-phe-leu-glu-gly-phe-val-arg-thr-ser-asn-ile-CTG-GGT-GGC-AAC-AAC-CTG-CCG-GCC-AAG-TTC-CTC-GAG-GGC-TTC-GTG-CGG-ACC-AGC-AAC-ATC-

AAG-TIC-CAA-GAC-GCC-TAC-AAC-GCC-GGT-GGG-CGC-CAC-AAC-GGC-GIG-TIC-GAC-TIC-CCG-GAClys-phe-gln-asp-ala-tyr-asn-ala-gly-gly- arg-his-asn-gly-val-phe-asp-phe-pro-asp-1074

ser-gly-thr-his-ser-trp-glu-tyr-trp-gly-ala-gln-leu-asn-ala-met-lys-pro-asp-leuagc-ggt-acg-cac-agc-tgg-gag-tac-tgg-ggc-gcg-cag-ctc-aac-gct-atg-aag-ccc-gac-ctgCAA-CG. -CAC-TGG-GTG-CCA-CGC-CCA-ACA-CCG-GGC-CCG- CCG-CAG-GGC-GCC-TAGCTCCGAACAGACA gln-arg- his-trp-val-pro-arg-pro-thr-pro-gly-pro-pro-gln-gly-ala-TER 1194

CAACATCTAGCNNCGGTGACCCTTGTGGNNCANATGTTTCCTAAATCCCGTCCCTAGCTCCCGCNGCNNCCGTGTGGTTA GCTACCTGACNNCATGGGTTT

MET-GLN-LEU-VAL-ASP-ARG-VAL-ARG-GLY-ALA-ACT-GCC-GGG-CCC-AGC-GCC-TGC-AGT-CTG-ACC-TAA-TTC-AGG-ATG-CGC-CCA-AAC-ATG-CAT-GGA-TGC-GTT-GAG-ATG-AGG-ATG-AGG-GAA-GCA-AGA-ATG-CAG-CTT-GTT-GAC-AGG-GTT-CGT-GGC-GCC-

 ${\it VAL-THR-GLY-MET-SER-ARG-ARG-LEU-VAL-VAL-GLY-ALA-VAL-GLY-ALA-ALA-LEU-VAL-SER-GLY-}$ GTC-ACG-GGT-ATG-TCG-CGT-CGA-CTC-GTG-GTC-GGG-GCC-GTC-GGC-GCG-GCC-CTA-GTG-TCG-GGT-

2

CTG-GTC-GGC-GCC-GTC-GGT-GGC-ACG-GCG-ACC-GCG-GGG-GCA-TTT-TCC-CGG-CCG-GGC-TTG-CCG-LEU-VAL-GLY-ALA-VAL-GLY-GLY-THR-ALA-THR-ALA-GLY-ALA-phe-ser-arg-pro-gly-leu-pro-181

7/60 GTG-GAG-TAC-CTG-CAG-GTG-CCG-TCG-CCG-TCG-ATG-GGC-CGT-GAC-ATC-AAG-GTC-CAA-TTC-CAAval-glu-tyr-leu-gln-val-pro-ser-pro-ser-met-gly-arg-asp-ile-lys-val-gln-phe-gln-241

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ser-gly-gly-ala-asn-ser-pro-ala-leu-tyr-leu-leu-asp-gly-leu-arg-ala-gln-asp-asp-AGT-GGT-GGT-GCC-AAC-TCG-CCC-GCC-CTG-TAC-CTG-CTC-GAC-GGC-CTG-CGC-ĠCG-CAG-GAC-GAC-301

phe-ser-gly-trp-asp-ile-asn-thr-pro-ala-phe-glu-trp-tyr-asp-gln-ser-gly-leu-ser-TTC-AGC-GGC-TGG-GAC-ATC-AAC-ACC-CCG-GCG-TTC-GAG-TGG-TAC-GAC-CAG-TCG-GGC-CTG-TCG-361

GTG-GTC-ATG-CCG-GTG-GGT-GGC-CAG-TCA-AGC-TTC-TAC-TCC-GAC-TGG-TAC-CAG-CCC-GCC-TGCval-val-met-pro-val-gly-gly-gln-ser-ser-phe-tyr-ser-asp-trp-tyr-gln-pro-ala-cys-421

GGC-AAG-GCC-GGT-TGC-CAG-ACT-TAC-AAG-TGG-GAG-ACC-TTC-CTG-ACC-AGC-GAG-CTG-CCG-GGGgly-lys-ala-gly-cys-gln-thr-tyr-lys-trp-glu-thr-phe-leu-thr-ser-glu-leu-pro-gly-

trp-leu-gln-ala-asn-arg-his-val-lys-pro-thr-gly-ser-ala-val-val-gly-leu-ser-met-IGG-CTG-CAG-GCC-AAC-AGG-CAC-GTC-AAG-CCC-ACC-GGA-AGC-GCC-GTC-GTC-GGT-CTT-TCG-ATG-

GCT-GCT-TCT-TCG-GCG-CTG-ACG-CTG-GCG-ATC-TAT-CAC-CCC-CAG-CAG-TTC-GTC-TAC-GCG-GGAala-ala-ser-ser-ala-leu-thr-leu-ala-ile-tyr-his-pro-gln-gln-phe-val-tyr-ala-gly-601

ala-met-ser-gly-leu-leu-asp-pro-ser-gln-ala-met-gly-pro-thr-leu-ile-gly-leu-ala-GCG-ATG-TCG-GGC-CTG-TTG-GAC-CCC-TCC-CAG-GCG-ATG-GGT-CCC-ACC-CTG-ATC-GGC-CTG-GCG-661

8/60 met-gly-asp-ala-gly-gly-tyr-lys-ala-ser-asp-met-trp-gly-pro-lys-glu-asp-pro-ala-ATG-GGT-GAC-GCT-GGC-GGC-TAC-AAG-GCC-TCC-GAC-ATG-TGG-GGC-CCG-AAG-GAG-GAC-CCG-GCG 721 168

TGG-CAG-CGC-AAC-GAC-CCG-CTG-TTG-AAC-GTC-GGG-AAG-CTG-ATC-GCC-AAC-AAC-ACC-CGC-GTC-188 787 SUBSTITUTE SHEET

trp-gln-arg-asn-asp-pro-leu-leu-asn-val-gly-lys-leu-ile-ala-asn-asn-thr-arg-val-

TGG-GTG-TAC-TGC-GGC-AAC-GGC-AAG-CCG-TCG-GAT-CTG-GGT-GGC-ÅAC-AAC-CTG-CCG-GCC-AAGtrp-val-tyr-cys-gly-asn-gly-lys-pro-ser-asp-leu-gly-gly-asn-asn-leu-pro-ala-lys-84.1

phe-leu-glu-gly-phe-val-arg-thr-ser-asn-ile-lys-phe-gln-asp-ala-tyr-asn-ala-gly-TTC-CTC-GAG-GGC-TTC-GTG-CGG-ACC-AGC-AAC-ATC-AAG-TTC-CAA-GAC-GCC-TAC-AAC-GCC-GGT-901

GGC-GGC-CAC-AAC-GGC-GTG-TTC-GAC-TTC-CCG-GAC-AGC-GGT-ACG-CAC-AGC-TGG-GAG-TAC-TGGgly-gly-his-asn-gly-val-phe-asp-phe-pro-asp-ser-gly-thr-his-ser-trp-glu-tyr-trp-961

Figure 5 (con't)

Ŧ 12

9/60

GG|C|-GCG-CAG-CTC-AAC-GCT-ATG-AAG-CCC-GAC-CTG-CAA-CGG-GCA-CTG-GGT-GCC-ACG-CCC-AACgly-ala-gln-leu-asn-ala-met-lys-pro-asp-leu-gln-arg-ala-leu-gly-ala-thr-pro-asn-268 1021

ACC-GGG-CCC-GCG-CCC-CAG-GGC-GCC-TAG-CTC-CGA-ACA-GAC-ACA-ACA-TCT-AGC-GGC-GGT-GAC-1081 288

thr-gly-pro-ala-pro-gln-gly-ala-TER

TGT-GGT-TAG-CTA-CCT-GAC-GGG-CTA-GGG-GTT-GGC-CGG-GGC-GGT-TGA-CGC-CGG-GTG-CCI-IGI-GGI-CGC-CGC-CGI-AGA-IGI-IIC-CIA-AAI-CCC-GIC-CCI-AGC-ICC-CGC-CGC-GGG-CCG-1141 1201 1261

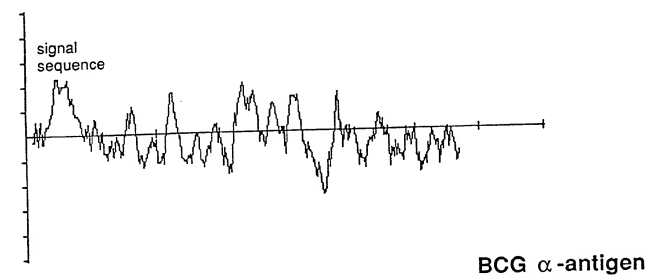
GCC-TAC-ACG-AAC-GGA-AGG-TGG-ACA-CAT-GAA-GGG-TCG-GTC

Figure 5 (con't)

Hydropathy

10/60

M. tuberculosis 32 kD protein



Hydropathy

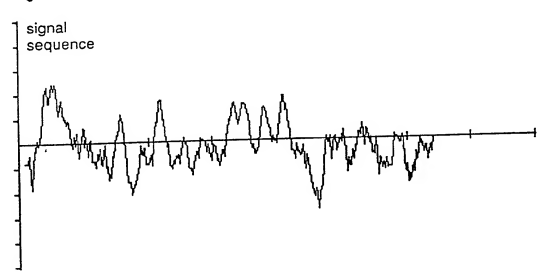


Fig. 6

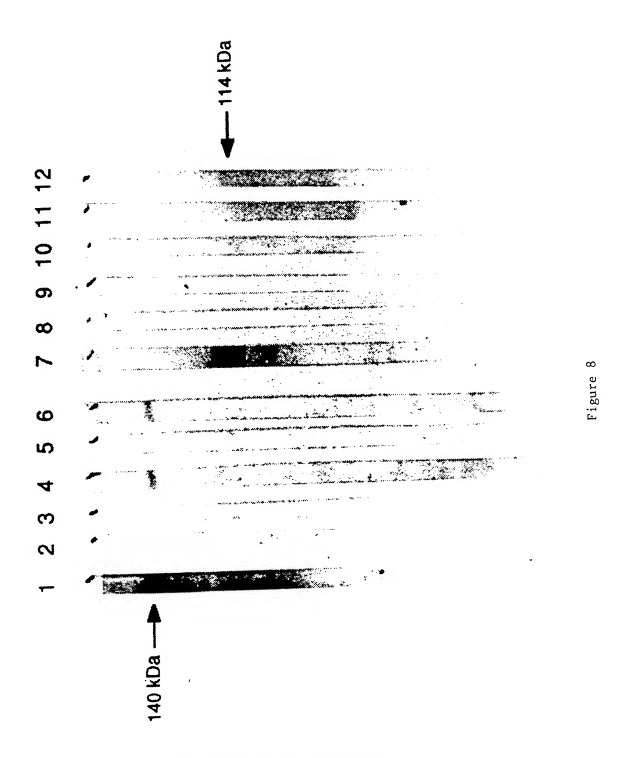
	11/6	o	
60 PSPSMGR :::::: PSPSMGR	120 :GQSSFYS :::::: :GQSSFYS	180 SALTLAIY ::::	240 IDPLLNVG ::: IDPTQQIP
50 PGLPVEYLQV ::::::::: PGLPVEYLQV 50	110 SGLSVVMPVG ::::::::: SGLSIVMPVG	170 18 SAVVGLSMAASSALTLA:::::::::::::::::::::::::::::::	230 PKEDPAWQRN ::::::::
40 GTATAGAESR ::::: GAATAGAESR 40	100 NTPAFEWYDQ ::::::::: NTPAFEWYYQ 100	160 HVKPTGSA ::::: :: AVKPTGSPSA 160	210 220 230 LIGLAMGDAGGYKASDMWGPKEDPAWQRNDP::::::::::::::::::::::::::::::::::::
SRRLVVGAVGAALVSGLVGAVGGTATAGAFSRPGLPVEYLQVPSPSMGR SRRLVNGAVGAALVSGLVGAVGGTATAGAFSRPGLPVEYLQVPSPSMGR GRRLMIGTAAAVVLPGLVGLAGGAATAGAFSRPGLPVEYLQVPSPSMGR 20 30 40 50	80 100 120 NSPALYLLDGLRAQDDFSGWDINTPAFEWYDQSGLSVVMPVGGQSSFY :::::::::::::::::::::::::::::::::::	140 150 160 CQTYKWETFLTSELPGWLQANRHVKPTG- ::::::::::::::::::::::::::::::::::::	200 220 240 SGLLDPSQAMGPTLIGLAMGDAGGYKASDMWGPKEDPAWQRNDPLLNVG :.:::::::::::::::::::::::::::::::::::
20 RLVVGAVGAA:::.::.:.::	80 PALYLLDGLR :::::::: PAVYLLDGLR 80	140 TYKWETELTS :::::::::: TYKWETLLTS	200 :LLDPSQAMGP :::::::: :LLDPSQGMG- 200
VDRVRGAVTGMSRRLVVGAVGAALVSGLVGAVGGTATAGAFSRPGLPVEYLQVPSPSMGR	70 80 100 120 DIKVQFQSGGANSPALYLLDGLRAQDDFSGWDINTPAFEWYDQSGLSVVMPVGGQSSFY; ::::::::::::::::::::::::::::::::::::	130 140 150 160 170 180 DWYQPACGKAGCQTYKWETFLTSELPGWLQANRHVKPTGSAVVGLSMAASSALTLAIY ::::::::::::::::::::::::::::::::::::	190 200 HPQQFVYAGAMSGLLDPSQAMG ::::::::::::::::::::::::::::::::::::
M. tub. BCG			

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300	HNGVFDFPD	::.:: HNAVENEPP	0						
290	QDAYNAGGG	SESSIBLE SES	290						
280	FVRTSNIKF	EVRSSNLKE	280						
270	NLPAKFLEG	NIPAEFLEN	270		GA	×	GA		
260	NGKP SDLGGN	.:::::::::::::::::::::::::::::::::::::	260	320	AMKPDLQRAL	X:::::::::::::::::::::::::::::::::::::	WGAQLNAMKGDLQSSLGA	320	
250	KLIANNTRVWVYCGNGKP SDLGGNNLPAKFLEGFVRTSNIKFQDAYNAGGGHNGVFDFPD	:::::::	250	310	SGTHSWEYWGAQLNAMKPDLQRALGA	••	WEYWGAQLN	310	
	KLIAN	KLVANNTR	240		SGTHS	••	NGTHSWEY	300	

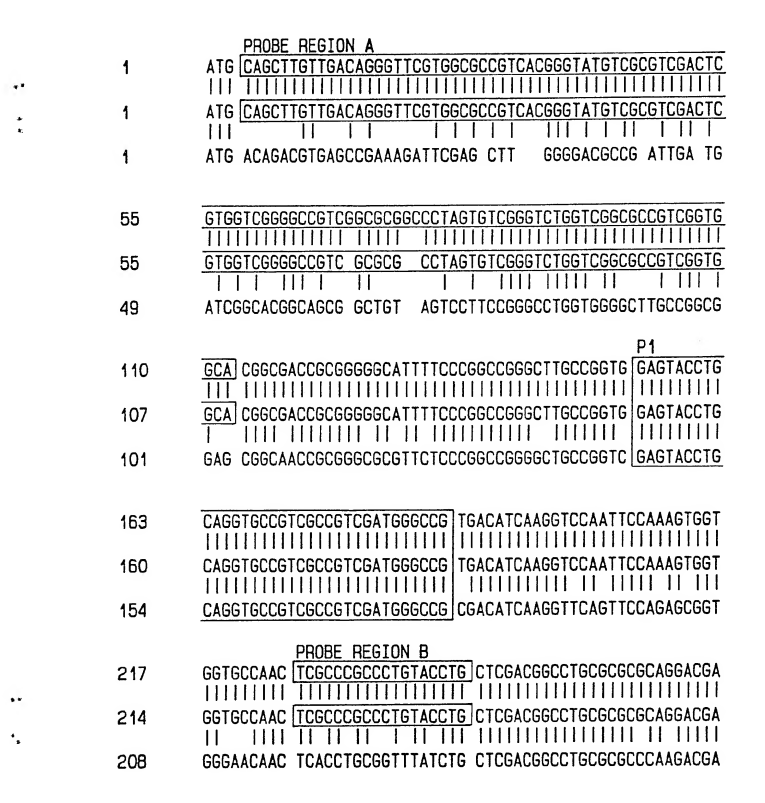
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14/60 Figure 9a



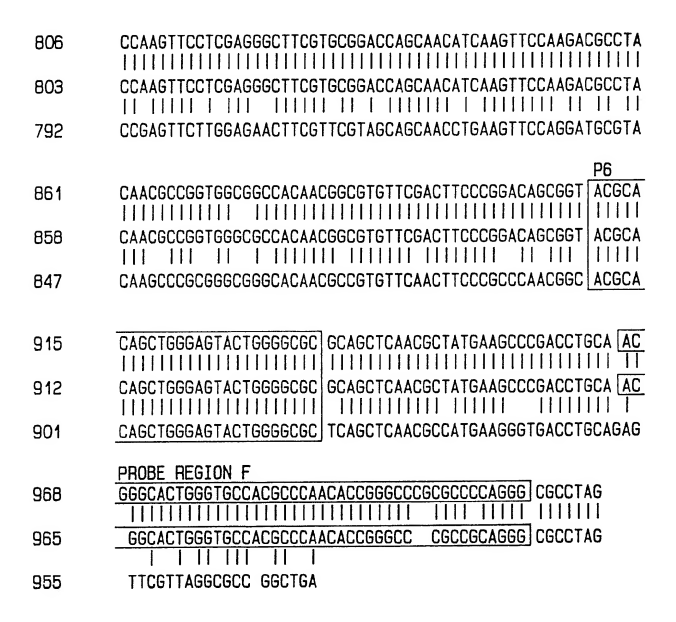
15/60 Figure 9b

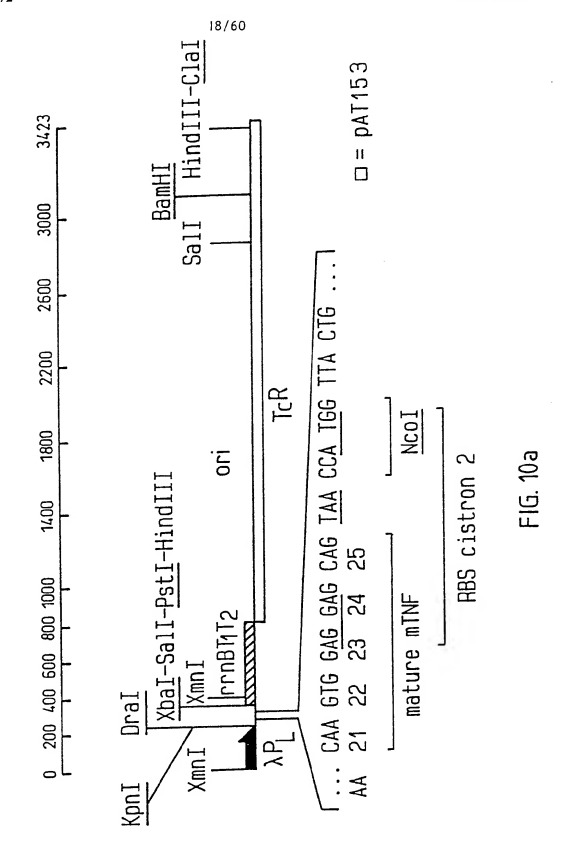
	P2
270	CTTCAGCGGCTGGGAC ATCAACACCCCGGCGTTCGAGTGGTAC GACCAGTCGG
267	CTTCAGCGGCTGGGAC ATCAACACCCCGGCGTTCGAGTGGTAC GACCAGTCGG
261	CTACAACGCTGGGAT ATCAACACCCCGGCGTTCGAGTGGTAC TACCAGTCGG
323	GCCTGTCGGTGGTCATGCCGGTGGGTGGCCAGTCAAGCTTCTACTCCGACTGGTA
320	GCCTGTCGGTGGTCATGCCGGTGGGTGGCCAGTCAAGCTTCTACTCCGACTGGTA
314	GACTGTCGATAGTCATGCCGGTCGGCGGCAGTCCAGCTTCTACAGCGACTGGTA
	P3 P4
378	CCAGCCGCCTGCGGCAAGGCCGGT TGCCAGACTTACAAGTGGGA GACCT TC
375	CCAGCCCGCCTGCCGCAAGGCCGGT TGCCAGACTTACAAGTGGGA GACCT TC
369	CAGCCCGGCCTGCGGTAAGGCTGGC TGCCAGACTTACAAGTGGGA AACCC TC
430	CTGACCAGCGAGCTGCCG GGGTGGCTGCAGGCCAACAGGCACGTCAAGCCCACC
427	CTGACCAGCGAGCTGCCG GGGTGGCTGCAGGCCAACAGGCACGTCAAGCCCACC
421	CTGACCAGCGAGCTGCCG CAATGGTTGTCCGCCAACAGGGCCGTGAAGCCCACC
	PROBE REGION C
484	GGAAGCGCCGTCGTCGGTCTTTCGATGGCTGCTTCTTCG GCGCTGACGCTGGCG
481	GGAAGCGCCGTCGTCGGTCTTTCGATGGCTGCTTCTTCG GCGCTGACGCTGGCG
475	GGCAGCGCTGCAATCGGCTTGTCGATGGCCGGCTCGTCG GCAATGATCTTGGCC

16/60 Figure 9c

	538	ATCTATC ACCCCCAGCAGTTCGTCTACGCGGGAGCGATGTCGGGCCTGTTGGAC
.	535	ATCTATC ACCCCAGCAGTTCGTCTACGCGGGAGCGATGTCGGGCCTGTTGGAC
	529	GCCTACC ACCCCAGCAGTTCATCTACGCCGGCTCGCTGTCGGCCCTGCTGGAC
		P5
	592	CCCTCCCAGGCGATGGGTCCCAC CCTGATCGGCCTGGCGATGGGTGACGC TGG
	589	CCCTCCCAGGCGATGGGTCCCAC CCTGATCGGCCTGGCGATGGGTGACGC TGG
	583	CCCTCTCAGGGGATGGG CCTGATCGGCCTCGCGATGGGTGACGC CGG
	645	CGGCTACAAGGCCTCCGACATGTGGGGCCCGAAGGAGGACCCGGCGTGGCAGCGC
	642	CGGCTACAAGGCCTCCGACATGTGGGGCCCGAAGGAGGACCCGGCGTGGCAGCGC
	631	CGGTTACAAGGCCGCAGACATGTGGGGTCCCTCGAGTGACCCGGCATGGGAGCGC
		PROBE REGION D
	700	AACGAC CCGCTGTTGAACGTCGGGAAG CTGATCGCCAACAACACCCGCGTCTG
	697	AACGAC CCGCTGTTGAACGTCGGGAAG CTGATCGCCAACAACACCCGCGTCTG
	686	AACGAC CCTACGCAGCAGATCCCCAAG CTGGTCGCAAACAACACCCGGCTATG
		PROBE REGION E
3	753	GGTGTACTGCGGCAACGGC AAGCCGTCGGATCTGGGTGGCAAC AACCTGCCGG
9	750	GGTGTACTGCGGCAACGGC AAGCCGTCGGATCTGGGTGGCAAC AACCTGCCGG
	739	GGTTTATTGCGGGAACGGC ACCCCGAACGAGTTGGGCGGTGCC AACATACCCG

17/60 Figure 9d Figure 9e





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From: PIGRI

			19	/60				
45	AAA TTT	AAA TTT	CTG	CTT	aaa Ttt	TAG	GGT	CCA
	AAA TTT	GAT CTA	ATA TAT	GCT	TTT AAA	CTG	CAT	TGC
39	TGC	GGT	GTG	GAC	AGG	AGC TCG	AAC TTG	ATC TAG
	999	TGC	393 939	GGT	AGG	ACA TGT	AGT TCA	TCA AGT
33	999	ATC T'AG	CTG	GAA CTT	ACC TGG	GTG	AGC	AGG TCC
	ATG	ACC TGG	CCA	CAT	GGT	CGA	AGG	CTG
27	ACA TGT	ATA TAT	ATA TAT	CAC	AGG TCC	ATT TAA	TGG	000 000
	CAA GTT	CAG	TAA	GAC	522 255	AAA TTT	AAG TTC	TCA
21	TAC	ATA TAT	ACA TGT	ACT TGA	AAG TTC	GTC	ACC TGG	AAC TTG
	ACC TGG	AAC.	TTG	ອວອ ວອວ	AAG TTC	GTA	ACC TGG	ACC TGG
15	CTC GAG	AAA TTT	GTG	GGA	CTG	CAA	CAA GTT	555 555
	TCT AGA	ATA TAT	000 000	GCA	550 000	GAT	TAG	AGG
9 -	GGA	CAT	CTG	TCA	TAA ATT	TAA	TCG	AGA TCT
	ວວອ ອອວ	ATT TAA	TCT	ACA TGT	AAT TTA	TGG	ACG	TGG
m -	TTC AAG	TAA ATT	TTA AAT	AGC	AAA TTT	TCA	999 ၁၁၁	TAC
	-	46	91	136	181	226	271	316

VT PA	GAT CTA GAA CTT TGG ACC GCG 09/07 CGC CGC CGC CGC CGC CGC CGC CGC								
G GAT C CTA	A GAA T CTT					AGG A	ນ ໓໓໓ ນນນ	CAA A GTT T	
ວອວ ອວອ	GCA	555 555 555	GTA GCAT	A ACT	990 S				
TTG	AAC TTG	090 909	550 005	GGA	TGG	AGT TCA	ວວອ ອອວ	CAT	
GTT CAA	CAG GTC	GTA	AAC TTG	TAG	GAC	CTG	CAA GTT	AGG	
GCT	AAT TTA	GCA	TGA	GAG	AAA TTT	CTC GAG	AAG TTC	990 229	
TTG	TTA	090 909	AAG TTC	CGA	TCG	GCT	ე <u>ნე</u>	ACT	
AGC TCG	AGA TCT	CTG	CAG	ATG	CAG	AAC TTG	GTT CAA	TAA	
(Con't) CCA AGC GGT TCG	TAC	TGC	ACT TGA	999	GCT	GTG	AAC TTG	CCA	
	TGA	ATT TAA	CGA	CTC	AAG TTC	TCG	TTG	555 555	
Fig. 10b TGC AGC ACG TCG	990 009	AGA TCT	TGC	GGT	CGA	TTG	GAT	525 252	
ACC TGG	TCA	AAC TTG	CCA	TGG	AAA TTT	TGT	ວອວ	GGA	
TCG	TTT AAA	TAA ATT	ACC	GTG	ATA TAT	TGT	GGA	GCA	
GAG	AGA TCT	TGA	CTG	GTA	CAA	ATC	255 522	ეე <u>ე</u>	
CTA GAT	AGA	GTC	CAC GTG	ATG TAC	CAT	TTT AAA	ე <u>ე</u> ნე	TGG	
AGT TCA	GAG	292 929	TCC	000 000	AGG TCC	CGT	AAT TTA	999 000	
361	406	451	496	541	586	631	919	721	
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225

TGC

GTT CAA

TTT AAA

TGG

225 552

CAG

TAC

525 252

CAC

AAC TTG

AAA TTT

CAA

AAA TTT

TGC

1036

						9 1	/60					
	CAA	GTT	CL	CGA	TCT	AGA	AAC ₉	TTG	TCA	AGT	GCT	
	CTA	GAT	TCC	AGG	GGA	CCI	CTT	GAA	AGA	\mathtt{TCT}	GCT	CGA
	TTT		GTA	CAT		TTT	TCC	AGG	AAA	TTT	TCT	AGA
	909	ວອວ	TAT	ATA	ATA	AGT TAT TAT TTT	AAA	GGT TTT	TAG	ATC	TAA	ATT
	TII		AAA	TTT	ATA	TAT	CCA	GGT	500	255	TGC GCG TAA	၁၅၁
	CLL		TTC	AAG	TCA	AGT	TGA	ACT	ACC	CIC IGG GGC	$\mathbb{T}G\mathbb{C}$	ACG
	299		ACA	IGT	GCT	CGA	rca	AGT			TTC	AAG
on't)	GAT		AAT	TTA	AAT	TTA CGA	ATC	TAG	CGT	GCA	TLL	AAA
Fig. 10b (Con't)	ACG	TGC	CTA	GAT	ATA	TAT	ATA	TAT	GAG	CIC	CTT	GAA
Fig.	CTG	GAC	TTT	AAA	CTG	GAC TAT	TTG	AAC	ACT	TGA	ATC	TAG
		TAG		TAA	ACC	TGG	TTT	AAA	TCC	AGG	GAG	CTC
	229			AAA	ATA	rat	TCC	AGG	CGT	GCA	CTT	
	AAG	TTC	TTG	AAC	ACA	CTC TGT	AGA	ACT TCT	TTT	AAA	CTT	GAA
	CAG	GTC	CTT	GAA	GAG	CTC	TGA	ACT	AGT	TCA	GAT	CTA
	AAG	TIC	ACT	TGA	CAT	GTA	AGG	TCC	GTG	CAC	991 AAG	TTC
	991		811		856		901		946		991	

	GCA	TAG	CTC CAG GAG	CGT	ອນອ ນອນ	TGG	ATT
	TCA	AGT TCA	TCG AGC	AGT TCA	AGG		CCG AAC TGA GAT ACC TAC AGC GTG AGC ATT GGC TTG ACT CTA TGG ATG TCG CAC TCG TAA
	GCT	CGT	ACC TGG	ATA TAT	ATA TAT	CCA	GTG
	CTG	AGC TCG	CAT	GTG GCG ATA AGT CAC CGC TAT TCA	ეე <u>ე</u> ეეე	AGC TCG	AGC
	TAA ATT	TGT ACA	CTA	GTG	TAC ATG	GCA CAC AGC CCA CGT GTG TCG GGT	TAC
	CGA AGG TAA GCT TCC ATT	TTC TAG TGT AGC CGT AAG ATC ACA TCG GCA	CAC CGC CTA CAT ACC TCG GTG GCG GAT GTA TGG AGC	CTG CCA GAC GGT	GAC GAT AGT TAC CGG ATA AGG CTG CTA TCA ATG GCC TAT TCC	GCA	ACC TGG
	CGA	TTC	CAC	CTG	GAT	CGT	GAT
on't)	CTC TTT TTC GAG AAA AAG	TCC AGG	CTG TAG GAC ATC	CTG	GAC	GTT CAA G	TGA
Fig. 10b (Con't)	TTT AAA	ATA CTG TCC TAT GAC AGG	CTG	CAG TGG CTG GTC ACC GAC	CAA	ນນນ ອອອ	AAC TTG
Fig. 1	CTC	ATA TAT	ACT	CAG	ACT (ວວອ ອອວ	
	CAA GTT	CAA	AGA TCT	TAC	TGG	GAA	ACA TGT
	TAC	TAC	TCA	TGT	GGT	GCT	CCT
	AGC TCG	AGA TCT	ACT TGA	TCC	095 000	ეე <u>ე</u> ეეე	CGA
	AAG TTC	505 050	ACC TGG	TAA ATT	TTA	GGT	GAA
	ATC	GAG	992 229	TGC ACG	GTC	AGC	AGC
	1081	1126	1171	1216	1261	1306	1351

ATC	CCA TAG	TTC	AAG	225	SSO	SGC	SOO		AGG
CCT	CCA	AGC	TCG		AAG		ລລລ	GGT	CCA
	\mathtt{TGT}	999	၁၁၁	GGT	CCA	CAG	GTC	TAC	ATG
550	၁၁၅	CGA	CCL		AGC	CGT	GCA	TTT	AAA
AGG	TCC	GCA	CGT	CTG	GAC	GCT	CGA	CCT	GGA
	CLL	AGC	TCG	GTC	CAG	GAT	CTA C	SSS	၁၁၅
GGA	CCT		CIC	ATA	TAT	TGI	ACA	ACG	$^{\mathrm{TGC}}$
AAG	TTC	CAG	GTC	TTT	AAA		AAA	GCA	CGT
မီ) (၄၄၄	299	GAA	CTT	ATC	TAG	GAT	CTA	CCA	GGT
Fig. 10b (Con't) TTC CCG AAG	AAG	TCG	AGC	\mathtt{GGT}	CCA	GTC	CAG	ACG	
JSJ	929	CCC		CCT	GGA	AGC	TCG	AAA	TLL
CCA	GGT	GCG GCA	CGT	ACG	IGC	TTG	AAC	GGA	CCI
GCG CCA	ວອວ	909	ນອນ	GAA	CTT	GAC		TAT	ATA
AAA	TTT	LAA	\TT	GGG	SSS	TCT	AGA	CCC	CGG
GAG	CIC	SSS	ນນຸນ	CAG	GTC	ACC	\mathtt{TGG}	GGA	CCL
1396 GAG AAA		1441 CGG 1		1486 CAG GGG		1531 ACC TCT		1576 GGA GCC TAT GGA	

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TAT ATA	CTG	TGA	TAC ATG	GCA	GGT	225 552
CGT	GAG	CAG	CTT GAA	GTC	ATC	AGC
CTG	AGT TCA	AGT TCA	AGA TCT	CAG	CGT GCA	CCT
rtc	FTG	CCG AGC GCA GCG AGT CAG GGC TCG CGT CGC TCA GTC	TCC AGA AGG TCT	GCT	CGC TCG CGT GCG AGC GCA	CAG
TCT AGA	CCT	GCA CGT	GTT	GTT CAA	9 29	525 252
TGT TCT T	CCG CCT 1	AGC TCG	525 252	GTT CAA	GTT CAA	555 ၁၁၁
ACA	TTA	299 922	GAC TTC CGC CTG AAG GCG	CAT	CAC GTG	GTA AGG CAA CCC CGC CAG CCT AGC CAT TCC GTT GGG GCG GTC GGA TCG
	GTA	CGA	GAC	ATT (TCG CTT AGC GAA	AGG
10b (Con't) TTG CTC AAC GAG	ACC TGG	GAA CTT	GCT CGA	ACC TGG	TCG	
Fig. CTT GAA	ata Tat	992 000	AGC TCG	AAG TTC	CAG	CCA
ອນນ ນອອ	TGG	GCA	AAG	ວອອ ອນນ	CAG	C TAA
GCT	CTG	990 ၁၁9	225 222	AAA TTT	CAG	TG
TTT AAA	ATT TAA	CTC	AAG TTC	555	TTG	TTC
CCT	CTG	ລອອ	AGG TCC	ACA TGT	GTT CAA	TCA
TGG	999 ၁၁၁	ATA TAT	ລອລ ອລອ	GAA	GAC	GAT
1621	1666	1711	1756	1801	1846	1891

24/60

252 525	ACA	AAT ⁵² TTA	999 ၁၁၁	ATA TAT	ວ <u>ອ</u> ອ ອວວ
			TGG	GGT	TCG
	GCA	GTG	AGG TCC	CAA GTT	TCC ATG TGC TCG CCG AGG TAC ACG AGC GGC
	TGC	GGA	TCG	AGA TCT	ATG
CTG	GTT CAA	CTT GAA	AGG TCC	922 299	TCC
000 000	TTG	ATT		GGA	CGT
GTG	222 555	CCA	CCA	ວວອ	CCA ACC CGT GGT TGG GCA
505 050	CAA GTT	GCT	CTT GAA		CCA
ອນອ ນອນ	TGC	TTG	ეე <u>ნ</u> ეეე		ATG
ATG	TTC	TGA	9 29		TCC
	ATG	AAT TTA	TGC		CAA GTT
555 555	GAT	AAG TTC	AGG TCC		CTA GAT
CTG	ATG	929 292	ე <u>ნე</u> ნენ		ອນອ
ACG	ე <u>ე</u> ეე ეეე	CTC	TTA AAT		505 005 000 050 550 555
CCA	GAC	GTT CAA			ວນນ ອອອ
1981	2026	2071	2116	2161	2206
	CCA ACG CTG CCC GGT TGC GAC GGG	CCA ACG CTG CCC GAG ATG CGC CGC GTG CGG CTG CTG GAG ATG GGT TGC GAC GAC GAC GAC GTC TAC GTG GTG TAC TAC GTG CTG GAC GAC GAC GAC GAC TAC TAC GTT TGC GTT TGC GTA TTC TGC CTA TAC ATG ATG ATG GTT CCC AAC CAA ACG CGT AAG	GAC GCG ATG CTC GAG ATG CGC CGC GTG CGG CTG CTG GAG ATG GGT TGC GAC GTC TAC GCG CTG CTG GAC GTC TAC TAC TTC TGC CAA GGG TTG GTT TGC GCA TTC TGC CTA AGG GTT CTT GGA GTG GTG CTG CTA AGG CTT CTC AAG AAG ACG GTT CCC AAC CAA ACG GTG GTG CAA GGG TTC CTA GGA GTG GTG CAA GAG GTG CTC CAA GGG TTC TTA ACT AAC CGA GGT TAA GAA CCT CAC CAC CAA GAG GCG TTC TTA ACT AAC CGA GGT TAA GAA CCT CAC CAC	GGT TGC GAG GTG CGC CGC GTG CGG CTG CTG GAG ATG GGT TGC GAG GTG CGG CAC GAC GAC GAC GAC GAC GAC TAC GAC GCG ATG GAT ATG TTC TGC CAA GGG TTG GTT TGC GCA TTC CTG CGC TAC CTA TAC AAG ACG GTT CCC AAC CAA ACG CGT AAG CAA GAG GCG TTC TTA ACT AAC CGA GGT TAA GAA CCT CAC CAC GGC ATT CTC CGC AGG TGC CGC CGG CTT CCA ATT CTT GGA GTG GTG CAA GAG GCG TTC TTA ACT CAC GAA GGT TAA GAA CCT CAC CAC GGC AAT CGC TCC ACG GCG CTT CCA ATT CTT GGA GTG TGG GGC AAT CGC TCC ACG GCG CTT CCA ATT CTT AGG TCC AGC TCC ACC	CCA ACG CTG CCC GAG ATG CGC CGC CTG CTG CTG CTG GAC ATG GGT TGC GGC CTG CTG CTG GAC CTC TAC GGG CTC TAC GCG CTC CAC GAC CTC TAC TAC CTA GCG CTT CCC AAC CAA ACG CTT TAC CTA TAC AAG ACG GTT CCC AAC CAA ACG CGT AAG CAA CCG TTC CAA GAG GCG TTC TTA ACT AAC CGA GGT TAA GAA CCT CAC CAC CAC CAC CAC CAC CAC CAC C

AAG TTC	GAT CTA	GCA [©] CGT	CCA GGT	CGT
TCG	CCT	30C	AGG	ე <u>ნე</u>
TGA	GTC	ACG	GGA	CCA
CAG	GCT	GCA ACG C	TGG	AGC
STC	GNA	CCT	TAA ATT	CGT
999 1	CTT GAA	TGG CCT (TCA	AGA TCT
TCA	GCG ATC CTT CGC TAG GAA	GCA	GAA	GCA
cga GCT	ე ეე ეეე	ACA TGT	GAA	CCA
10b (con't) 3 TGA CGA 2 ACT GCT	CGA	TGG	CGA	ACG
Fig. 1 CCG GGC	ე <u>ე</u> ნე	ອອວ ວວອ	CGG AAG GCC TTC	CGA
TCG	GAG	CCT	000 000	TCG
AAA TTT	TAA ATT	CTA	ອນອ ນອນ	ນນນ
CAT	TGG	CAT	TGC	TCC AGC CTC GCG AGG TCG GAG CGC
225 862	5 22	CGT	CGA	AGC
AGG	TTA	GGT	TCC	TCC
2251	2296	2341	2386	2431

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	2476	2521	2566	2611	2656	2701	2746
	ეე <u>ნ</u> ეეე	TGG	TTC	GAA	GTC	CGA	AGG
	ວອອ	TGG	CGA	AGC	CTA	TAG	CTC
	CCA	၁ 25	ATA TAT	GGT	CGA	TCA	TCA
	TGC	GAC	ວອອ ອວວ	CCT	GTT	TGC	AGG
	225 252	CAG	CAA GTT	ອນອ ນອນ	GCA TGA CGT ACT	<u>ອ</u> ອອ	GCA
Fig.	CGA	TGA	252 525	CGA GCT	TGA	090 909	TCG
201	CGA TAA TGG GCT ATT ACC	CGA	ACA TGT	AAA TTT	TAA	555 ၁၁၁	TCG GTC AGC CAG
		AGG TCC	922 299	TGA	AGA TCT	ACC	GAC
	CCT GCT GGA A	CGA AGG CTT GCT TCC GAA	GGC CGA TCA TCG TCG CGC CCG GCT AGT AGC AGC GCG	TGA CCC AGA ACT GGG TCT	AGA AGA CAG TCA TAA TCT TCT GTC AGT ATT	ACC GGA AGG AGC TGA TGG CCT TCC TCG ACT	GAC GCT CTC CCT CTG CGA GAG GGA
	GCT	GAG	TCA	AGA TCT	CAG	AGG	CTC
	rct aga	CGA	TCG	ວອວ ອວອ	TCA	AGC	CCT
	505 050	GGG CGT	TCG	CTG	TAA	TGA	TAT
	CGA GCT	CGT GCA	929 292	ວອອ ອນວ	GTG	CTG	TAT GCG ACT ATA CGC TGA
	AAC TTG	GCA	TCC	GCA	000 000	GGT	ACT TGA
	GTT CAA	AGA TCT	AGC	27/60 ECH ECH ECH ECH ECH ECH ECH ECH ECH ECH	CGA	TGA	CCT

TGG

GTG

225 252

GGA

ACA TGT

TCC

GGA

AGA

CGT

225 252

GTC

TGC

CGA

CCA

 \mathcal{CGG}

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	G
	TGA
	CGT
n't)	GTA
Fig. 10b (Con't)	GTA
Fig.	CCA
	AGC CCA
	AGC
	GGA
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255 522	GTC	929 292	TGT 628/82 ACA ACA	TGC
GCA	ACA TGT	AAG TTC	TGA	TGA
TGA	CCA	AAC TTG	ეე <u>ნ</u> ენე	ეე <u>ე</u> ეეე
CGT	909 090	CGA	CAT	505 050
500	TGG	505 050	999	TGG
TGA	AGA TCT	CCA	CTT	CTG
GGT	AGG TCC	TAC	GAT	CAC
GTA	GCA	CCA	999	ე <u>ე</u> ნე
GTA	CAT	CCA	GAG	CAA GTT
CCA	GTG	CTG	5 22	CAG
AGC	ATG TAC	922 255	AGT	909 090
	GGA	000 000	CGA	AGG TCC
GGA AGC CCT TCG	CAA	CCA	990	TAT
TTA	ე <u>ე</u> ნე	000 000	TGA	CGA
GCA	ე <u>ე</u> ნე	999 ၁၁၁	TCA	ეე <u>ნ</u> ეეე
2791	2836 CCG GGC GGC	2881	2926	29.71

CGA AAG TTC 252 GTA CAA CTC TGG TAG CGA AGT CGT TCG TGA CCA TCG 3061

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GAA CTT	GCA CGT	GCA ² CGT	CAT	ATT TAA
CGA	GCA	999 ၁၁၁	CAG	TAG
CTC GAG	CTA	TAT ATA	CTA	TGT
GTG	252 525	CGA	TGC	CAT
ACA TGT	ATA TAT	GGA	CTA GAT	ე ეე ე
225 552	CAT	AAT TTA	AGC TCG	TGA
GGT	ACG	ນນອ	CCA GGT	CGA
AGC TCG	TCA AGT	TGT ACA	TAA ATT	TGA
CMA	GCA	TGC	GCA	GGA
522 255	ATT TAA	CGA	255 522	CGA
522 255	GAA	TGG	GTA	TGC
522 255	ATA TAT	GAC	GCA	999 999
CTG GGC GAC CCG	ອວອ	AGT TCA	ე <u>ე</u> ნე	TGA
SGA	GTG	CAT AGT GTA TCA	299 922 922 299	999 300
GCA	225 552	ອນອ ນອນ	AGA TCT	CCA
3106 GCA C CGT (3151 CGG GTG CGC GCC CAC GCG	3196 CGC (3241	3286 CCA GGG TGA GGT CCC ACT
(.)		• •		

			30/60
AAC TTG	AAT TTA		T;
ATA TAT	GAG CTC		702 T;
GTG ATA Z	CAT		
CT (LAA (FTT (967 G;
TA P	TC A		96
AAT TTA ACT TTA AAT TGA	GCT GTC AAA CGA CAG TTT		915 C;
GC A	TAA G ATT C		915
Fig. 10b (Con't) TGC GTT AGC P ACG CAA TCG I	TGA T ACT A		A;
3. 10b	CGA T		9
Fig. C TC	T. C.(3423.
GAC	TAT ATA		
CCT	GCT		es is: ition: 0060.
GTG	AAA TTT		of bases is: composition:
ACG	ATT		of con
TAC ATG	ງຄວອ		mber ence
TCA	TAC	TAA ATT	Total number of base DNA sequence composi Sequence name: NIPS
3331 TCA TAC ACG GTG AGT ATG TGC CAC	3376 TAC CGC ATT AAA ATG GCG TAA TTT	3421 TAA ATT	Tota DNA Sequ

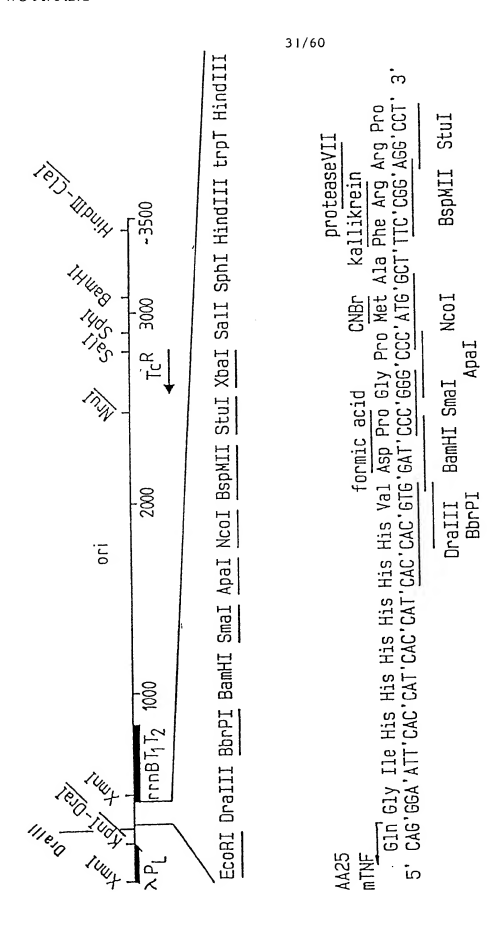


fig.116

				22/4	· 0				
	45	AAA TTT	ATA TAT	TAC ATGE	CTC	TTA	TGT ACA	TCA	GAG
		GCA	GTG	TGA	ACG	GGT	ອອລ ລລອ	AAT TTA	ລອອ ອວລ
	39	CCT (252 525	ລວອ	GTG	GGA	CAA GTT	000 000	TTT AAA
) 555 555	TCT (TGG	AAG TTC	CCA GGT	TGA	GCA	522 255
	33	TGC (CCA '	CAC	ATG	GTA	GAG	GGA	CAT
		CAA GET	TAA (ATT	TAC	ACC	ນນນ	TTC	GGA	550 225
	27	AAA (TTT (AGA	AAA TTT	ACC	GCA	AAA TTT	AGT	000 000
F1g. 11b		ACC TIGG	TAC	CAT	CTG	AGG	TCA	CCA	TCC
Ξ	21	CCT	ACA	TGA	GCA	AGA TCT	TAG	CCA	GGA
		TCA	AAA TTT	TGT	GAC	TGA ACT	AAG TTC	AAA TTT	CGT
	15	CTC	TAA	225 552	CAG	999 ၁၁၁	ATC TAG	AGC	CCA
т		GNT	ATA TAT	TGG	CAG	AAG TTC	AAG TTC	CGT	TCA
pmTNF JPH	6-	222	TTC	CTC	CAT	ATT TAA	GGT	CGT	CCA
pmTN		TCC	AAA TTT	TAT ATA	GCA	AAA TTT	CAT	CCA	TCA
From:	ო -	AAT	AAT TTA	AAT TTA	TGA	TTA	AAT TTA	AGC TCG	CCA
Fr		H	46	91	136	181	226	271	316

ບ ຫ	T &	K H	ပ္ ည ³	ACG _{9/8} TGC	<u>ن</u> ي	T.	AS T	AC FG
ອອນ ລວອ	TTT AAA	AGA			AGG.	ACT	TGA ACT	AAC
ອອວ ລລອ	CTG	ATC TAG	CAG	GAA	AGT	AAG TTC	TCC	AGC TCG
TAA ATT	TGG	TAA ATT	000 000	AGT TCA	GAG	CGA	CTC	CGA
AAG TTC	GCT TGG CGA ACC	GAT	TGG	AGA TCT	TGC	AGT	ACG	TTG
AGT TCA	CAA GTT	ACA TGT	992 229	CTC	CCA	CTC	TGA	ACG
TTA AAT	550 225	GAT	TTT AAA	GAA	TCC	AGG	000 000	TGA
	GAT	CCT	GAA	992 229	GTC	GAA	TGT	ATT
GCA AGC CGT TCG	NTT NAA	CAG	ACA TGT	CAT	ລລວ	AAC TTG	GTT	550
Fig. CAT GTA	TTN	TTT AAA	AAA TTT	999 ၁၁၁	TGT	TAA	GTT	GAG
ວນອ ອອນ	ATT TAA	GAT	GAT	TGA	TAG	AAA TTT	TCT	000 000
GAC	ອນນ	GAA	TCT	ACC TGG	TGG	ATC	TTA	525 252
GTC	500 255	AGA TCT	000 000	555 ၁၁၁	CGA	500 055	GTT	ATC
AGA TCT	GCT	ATG	AAG TTC	GGT	<u>ნენ</u> ენე	CCA	TTC	CAA
TCT AGA	TCC AGG	225 252	CAG	GGT	TAG	CTG	CCT	GGA
550 225	AGT TCA	TGG	ACG	5 25 25 25 25 25	ე <u>ე</u> ნე	GAA	222 888	GTA
361	406	451	496	541	586	631	919	721

CGT CGTGCA CAA CGT AAT TTA TAA ATT GCA CCA GGT AAC 252 266 CCCGAC GGG TCA AGT GTTCAA CILG AAA TCT AGA AGA TCT GTA 229SSO CAT GTA GAA CAT CTT AAA TTT AAA GTC ATA TAT TAC AGA ATG TCT ATG TAC CAT GTA TTT AGC TAA TAA TAA 550 505 000 CCC TCC CTG TGA TGA ACT TTCAAG CGGTGA Fig. 11b (Con't) AGA TCT CCA TCC TTT TTT 999 222 CTG TTG GTT CCT TTAATT CCA TAA TTC TTC GAT CAA TGT AGG GTT 222 GTT CAA GAA TTT AGA AGA TCT TCT 000 000 AGG TGA ATG GGT AGC GAG GGTCCA CAA ACG TTA AAATTT GAG GAT CIC CTA GAG TTA GAT CTA TAC GAT CTA SCC CGC TTT CGC SCG MAA AAG TTC AAA CCC 555 TAT ATA AAG TAG ATC CCC 1036 976 991 856 901 991

272						101,21
TTG	TGG	225 225	ATA S TAT	CGA GCT	GGA	992 229
GGT	AAC TTG	GTA	TAC	TGG	ACC TGG	ACA TGT
GGT	GGT		990	CAG	GTT CAA	CAC ACA GTG TGT
	GAA CTT	TGT CCT TCT AGT ACA GGA AGA TCA	CTC TGT AGC ACC GCC GAG ACA TCG TGG CGG	ACC AGT GGC TGC TGC CAG TGG TCA CCG ACG ACG GTC	CTC AAG ACG ATA GAG TTC TGC TAT	STG
ACC T		CCT	AGC	TGC	ACG	TTC
GCT ACC AGC CGA TGG TCG	TTT	TGT ACA	TGT	522	AAG TTC	GGG TTC C
	rcr AGA	TAC	CTC GAG	AGT TCA	CTC	000 000
11b (con't) ACC ACC TGG TGG	AAC TCT TTT TCC TTG AGA AAA AGG	aaa TTT	GAA CTT	ACC	GGA	AAC GGG (TIG CCC (
Fig. 1 AAA TTT	ACC	ACC	CAA	GTT CAA	GTT CAA	CTG
AAA	GCT	GAT	CTT GAA	CCT	ວວອ ອອວ	ວວວ ອອອ
AAC	AGA	GCA	CCA GGT	AAT TTA	TAC	GTC
GCA	TCA	AGC TCG	CCA	GCT	TCT	ე <u>ნე</u>
CTT	GGA	CAG	AGG TCC	TCT AGA	GTG	GCA
CTG	550 000	CAG	GTT	505 000	GTC	500 055
CTG	TTT AAA	CTT GAA	GTA	CCT	TAA ATT	TAA
1081	1126	1171	1216	1261	1306	1351

1't)
(Con
11b
Fig.

	GGA	36/6 CC CC CC CC	GNG	CIC	990	၁၁၅	GTC	CAG
T.C.T.	CCC	ອນນ	CAC	GTG	TGT	ACA	CTC	GAG
GGA	AAA	TTT	BCG	CGC	TCC	AGG	ATG	TAC
TAT.	GAG	CIC	AGA	TCT	TAG	ATC	GTG	CAC
	AGG	ICC	AGG	ICC	TTA	AAT	TTT	AAA
TGA	CGA	CCL	AAC	TTG	TCT	AGA	ATT	TAA
C.T.	TCC	AGG	SSS	ນນອ	GTA	CAT	\mathtt{TCG}	AGC
$G.\Gamma.G$	GCT	CGA	GGT	CCA	CTG	GAC	909	U U U
GAT.	CAC	GTG	CAG	GIC			TGA	ACT
C.I.G	CGC	909	SSS	ລລອ	AAA	TTT	ACT	TGA
TTG	AAG	TTC	AAG	TIC			CTG	GAC
ວ <u>ອ</u> ວ	AGA	TCT	GGT	CCA	AGG	TCC	CCT	GGA
L CCI C	TIG	AAC	TCC	AGG	TCC	AGG	CCA	CAA AGC GGT GGA
GAA	GCA	CGT	GTA	CAT	GCT	CGA	TCG	AGC
GTC	TGA	ACT	CAG	GIC	GGA	CCT	GTT	CAA
	1441		1486		1531		1576	
	CTG GAT GTG GCT TGA CTC TAT GGA	CCT CGC TTG CTG GAT GTG GCT TGA CTC TAT GGA TTG AGA AAG CGC CAC GCT TCC CGA AGG GAG AAA	TTG AGA AAG CGC CAC GCT TCC CGA AGG GAG AAAAAC TCT TTC GCG GTG CGA AGG GCT TCC CTT TTT	TTG AGA AAG CGC CAC GCT TCC CGA AGG GAG AAA AAC TCT TTC GCG GTG CGA AGG GCT TCC CTC TTT TCC GCG GTG CGA AGG GCT TCC CTC TTT TCC GGG CAG GGT CGG AAC AGG AGA GCG	TTG AGA AAG CGC CAC GCT TCC CGA AGG GAG AAA GGC AAC TCT TTC CGG GTG CGA AGG GCT TCC CTC TTT CCG TCC GGT AAG CGG CAG GGT CGG AAC AGG AGA GCG CAC AGG CCA TTC GCG CAC GTC CTC TTT CGG CAC GTC CTC TTG TCC TCT CGC CAC	TTG AGA AAG CGC CAC GCT TCC CGA AGG GAG AAA AAC TCT TTC GCG GTG CGA AGG GCT TCC CTC TTT TCC GGT AAG CGG CAG GGT CGG AAC AGG AGA GCG AGG CCA TTC GCC GTC CCA GCC TTG TCC TCT CGC TCC AGG GGG AAA CGC CTG GTA TCT TTA TAG TCC	TTG AGA AAG CGC CAC GCT TCC CGA AGG GAG AAA AAC TCT TTC GCG GTG CGA AGG GCT TCC CTC TTT TCC GGT AAG CGG CAG GGT CGG AAC AGG AGA GCG AGG CCA TTC GCC GTC CCA GCC TTG TCC TCT CGC AGG CCA TTC GCC GTC CCA GCC TTG TCC TCT CGC AGG CCC TTT GCC GTC CAT AGA AAT ATC AGG	TTG AGA AAG CGC CAC GCT TCC CGA AGG GAG AAA AAC TCT TTC GCG GTG CGA AGG GCT TCC CTC TTT TCC GGT AAG CGG CAG GGT CGG AAC AGG AGA GCG AGG CCA TTC GCC GTC CCA GCC TTG TCC TCT CGC AGG CCA TTC GCC GTC CCA GCC TTG TCC TCT CGC CCA CTC TTT GCC GTC CTG GTA TCT TTA TAG TCC AGG TCC CCC TTT GCG GAC CAT AGA AAT ATC AGG CCA CCT CTG ACT TGA GCG TCG ATT TTT GTG ATG

		3 /	//60			
TTT AAA	TCC	TGA	CGA	CCA	CTC	525 252
CTT GAA	CTT	CTT GAA	CAG	TTT AAA	TTG	GCT
		525 252				TTC
	TCA	TAT ATA		ACT TGA	TTC	TTC ACG
CAG	TGC	255		CTG	CCA	ອນອ
			ລອອ			AGT
						AGC TCG
		TGT ACA		GGA		AGC AGC TCG TCG
ATG TAC						TGC
					CAC	TTT AAA
						ACG TGC
						CAG
						TCG AGC
		TGC		GTC	GAC	AGG TCC
1621	1666	1711	1756	1801	1846	1891
	AGG GGG GCG GAG CCT ATG GAA AAA CGC CAG CAA CGC GGC CTT TCC CCC CGC CTC GGA TAC CTT TTT GCG GTC GTT GCG CCG GAA	AGG GGG GCG GAG CCT ATG GAA AAA CGC CAG CAA CGC GGC CTT TCC CCC CGC CTC GTT GCG CTC GTT GCG GAA ACG GTT GCG GTT GCG GAA GAA AAC GAC CGG AAA ACG AGT GTT CTT TGC CAA GGA CCG GAA AAC GAC CGG AAA ACG AGT GTA CAA GAA	AGG GGG GCG GAG CCT ATG GAA AAA CGC CAG CAA CGC GGC CTT TTT ACG GTT CCT GGA TAC CTT TTT GCG GTC GTT GCG CCG GAA AAA ACG GTT CCT GGC CTT TTG CTG GCC TTT TGC TCA CAT GTT CTT TCC TGC CAA GGA CCG GAA AAC GAC CGG AAA ACG AGT GTA CAA GAA AGG TGC GTT ATC CCC TGA TTC TGT GGA TAA CCG TAT TAC CGC CTT TGA ACG CAA TAG GGG ACT AAG ACA CCT ATT GGC ATA ATG GCG GAA ACT	ACG GGG CCG CTC GGA TAC CTT TTT GCG GTC GTT GCG GCC CTT TTT GCG GTC GTT GCG CCG GAA AAA ACG GTT GCG GTT GTT TCC CTT TTG CTG GCC TTT TGC TCT GTT CTT TCC TTT TGC TCT GTA GTA ACG ACT GTA CTA GTA ACG ACT GTA CTA GAA ACG GAA AAC GAC CGG AAA ACG AGT GTA CTA TAC TCT TCC TGT GCC TAT TAC CCG TAT TAC CGC TAT TCC TGT GCC TAT TAC CGC TAT TAC CGC TAT TAC CCC TCT TCC TCT ATC CCC TCT ATC GCC TCT ATT GCC TTT TCC CTT TC	ACG GGG GCG GAG CCT ATG GAA AAA CGC CAG CAA CGC GGC CTT TCC CCC CGC CTC GTT TTT GCG GTC GTT GCG CCG GAA ACG GTC CTT TTT GCG GTC GTT GCG CCG GAA ACG GTC CTT TG CTG GCC TTT TGC TCT GTT GCG CTT TGC CTT	AGG GGG GTT TTT GGG CTG GGG CTT TTT TTT GGG GTT GTT TTT ACG GTT GTG GTT TTG GTG GTT GTG GTT TTG ACG GTT TTG GTG GTG

37/60

GAC

GCA

GAG

222

505 000

CAA

ACG

252

ACC

TGC

CCA

GCT

255

522 255

GGT

2206

				-							
	AGC	GTG	CAC	\mathtt{TGG}	ACC	8/69 CC CC 8	ວອວ	GAG	CIC	CGA	GCT
	550 225	ລວລ	999	\mathtt{TGC}	ACG	TTT	AAA	TTG	AAC	GGT	CCN
	555 000	GCA	CGT	SGC	ອນນ	TGG	ACC AAA	TTC	AAG	TCA GGT	AGT
	GGC AAC	TGC	AGT ACG	TGC	ACG	GGT	CCA	CAA	GTT	CAT	GTA
	500 060	TCA	AGT	SCG	ລອລ	AAG	TTC	CIC	GAG	TTC	AAG
	TAA	CGA	GCT	CCC	ອອວ	CCC	550	TGG	ACC	299	ອນນ
t)	GCT AAC CAG	GCA	CGT	\mathtt{TGC}	ACG GGC TCT ACG CGG CGC ACG ACG	TCT	ACC TAT ACA AGA CGG TTC CCA	GAT	TCT TAA CTA	GCC GCC GGC TTC CAT	SSS
Fig. 11b (Con't)	AAC	GGA	TGC TGT CCT CGT	AGA	TCT	TGT	ACA	ATT	TAA	225	990
Fig. 11	GCT	ACA	TGT	SCG	299	ATA	TAT	AGA	TCT	GGT	CCA
	TCT	ACG	TGC	\mathbb{L} CC	ACG			GCA	AGG CGT	CGA	GCT
	CAT	TCA	AGT	၁၅၁	ອວອ	CGA	GCT	TCC	AGG	TAG	ATC
	ATT	TCC	AGG	CAA	GTT	ACG	TGC	TTC	AAG	CGT	GCA
	GTG ATT	999	၁၁၁	ACC	\mathtt{TGG}	550	225	CAG	GIC	TGA ATC	TAG
	TCG	225		AGG	TCC	$\mathbb{T}GG$	ACC	TCA	AGT	TGA	ACT
	GTA	CTA	GAT	CCC	CGG	AGA	TCT	CAT	GTA	TGG	ACC
	1936	1981		2026		2071		2116		2161	

Fig. 11b (Con't)

TGT	ACA	AGT	TCA 139/	CIGS	GAC	CAA	GTT	CCC	
CCA	GGT	TCC	AGG	AAG	m TTC	CTG	GAC	AAT	
GTT	CAA	550	ວວອ		AAC		ອວວ		GTA
	999		GIC	TCC	AGG	CAT	GTA	AAT	TTA
CAA	GTT	GAT	CTA	CGA	CCL	CAG	GTC	AAG	TIC
	ACG		CTG	GAG	CIC		CCT	GAG	CIC
	GGT	CGT	GCA	CCC	ອວອ	CCT	GGA		TCG
AAT	TTA	၁၅၁	929	AGC	TCG	CTG	GAC	GGA	
TAC	, ATG	AAT	TTA	AAG	TTC	TAC	ATG		CGG
225	990	ATA	TAT		CCA	ATC	CAG TAG		990
	ອນນ		ರಿ ಬ	GCT	CGA	GTC	CAG	GAT	CTA
299	ರಿ	ggc	ອວວ	TAG	ATC	GIC	CAG	CCC	SSS
TAG	ATC CCG	CGA	GCT	AGT	TCA	ATG	TAC	CAT	GTA
GTA	CAT	CGC	3CG	CGA	GCT TCA A	CTG	GAC	GGG	CCC
AAG	TIC	GCT	CGA	GAT	CTA	TCC	AGG	ນອນ	SCG
2251		2296	CGA (2341 GAT		2386	AGG GAC TAC CAG	2431	GCG CCC GTA

GAC

GCT

GGA

GAA

299 522

CCA

525 252

255 222

990 000

CAT

AGT

GAT

GAC

522 255

TGC

2746

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11b (
Fig.

	550 000	990 009	⁴ 522 255	ອວອ ວອວ	TGC	AAG TTC
	GTA CAT	CTC	GAG	CGT	ອນອ ນອນ	CAT
	GAC GTA CTG CAT	TTT	AGC	CAT	GAG	AGT
	CAA	GGC CTG C	GAA GGC TTG AGC CTT CCG AAC TCG	TAC CGC AAG CGA CAG GCC GAT CAT ATG GCG TTC GCT GTC CGG CTA GTA	CTC GCC GAA AAT GAC CCA GAG CGC GAG CGG CTT TTA CTG GGT CTC GCG	TTG CAT GAT AAA GAA GAC AGT AAC GTA CTA TTT CTT CTG TCA
	CAG	5 22	ອນນ ນອອ	<u>ອ</u> ອວ ວວອ	GAC	GAA
	CGC CYG C	AAT TTA	GAA	CAG	AAT TTA	AAA TTT
't)	CGC GAA GCG CTT	GCC GGC GAT CGG CCG CTA	GGC GGG ACC AGT GAC	CGA	GAA	GAT
11b (Con't)	929 292	500 005	AGT TCA	AAG TTC	550 225	CAT
Fig. 1	CGT	550 009	ACC TGG	525 252	CTC	TTG
	TCG	CAT)))	TAC	GCG GTC C	GAG
	550 225	525 252	522 255	GAA	ე <u>ნე</u> ნენ	TAC (
	CCA	522	GGT	TCC	AAA TTT	TCC
	CAT	GTC	TTT AAA	GAT	353	CTG
	500 000	505 050	ACG TTT TGC AAA	CAA	CCA	CAC
	GAA	CAG	GAA	GTG	GCT	225 552
	2476	2521	2566	2611	2656	2701

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Con	
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Fig.	
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	ATG	GTT CAA	ອອວ ວວອ	GAAA CTTO	CCC ATC GGG TAG	ອອນ ນນອ	GAC
	CTT GAA	550 205	522 255	990	999 ၁၁၁	ອນນ ນອອ	CCA CAG GGT GTC
	TCC AGG	GAG	GAT	CAC	TTC	TGT ACA	CCA
	CTC TCC GAG AGG	GTT	GGA	GGG GCC TGC CAC CAT ACC CAC GCC CCC CGG ACG GTG GTA TGG GTG CGG	GTG GCG AGC CCG ATC TTC CAC CGC TCG GGC TAG AAG	GCC AGC AAC CGC ACC CGG TCG TTG GCG TGG	GCG TCC GGC GTA GAG GAT CGC AGG CCG CAT CTC CTA
	ACG	TAG	CAA	CAT	299 922	505 050	GAG
	TCG	TAG	ATG	CAC	AGC	AAC TTG	GTA
	CGG TCG ACG GCC AGC TGC	GCC CAG TAG CGG GTC ATC	TGG TGC ATG ACC ACG TAC	TGC	ລອວ	AGC TCG	500 055
<u>.</u>	CAT	990 229	TGG	55 0	GTG	552 225	TCC
LIE TID (CON C)	GGG CAT	GCA	GAA	ეეე ეეე	GAA	ອນນ ນອອ	000 000
. 18.	CAA	GAA	AAG TTC	CAC	555 222	ATA	GAT
	TCT	TAG	505	522 255	GAG	GAT	CAC GTG
	900 099	CAT GTA	ອນອ ນອນ	999 ၁၁၁	CAT	ອນນ	500 055
	GNA	CTG	ອນອ ນອນ	TCC	GCT	GTC	<u>ულე</u> ეეუ
		CTC	CAC	CAG	AGC TCG	GAT	GAT
	TGG GTT ACC CAA	CGA	GAG	CAA GTT	ACA	GGT	GGT
	2791	2836	2881	2926	2971	3016	3061

	R	
	C GTA GTC GAT AGT GGC TCC A	
	299	
	AGT	
	GAT	
	GIC	
on't)	GTA	,
Fig. 11b (Con't)	CGC	
Fig.	GAT	i
	GGT CGC CAT GAT CGC	
	CGC	
	GGT	1
	TGT	1
	SSS	1

	4.3	/60		
TAG	TGC	/60 DDD	GAT	555 555
AAG TTC	CAG	TAG	GAC	TAT ATA
TCC	GGA	ATA TAT	ATG TAC	GCC TAT CGG ATA
ອນນ ນອອ	GCG GTC CGC CAG	CAT CAA CGC ATA TAG GTA GTT GCG TAT ATC	GGC GAT GCT GTC GGA ATG CCG CTA CGA CAG CCT TAC	CGG CAT AAC CAA GCC GTA TTG GTT
AGT ICA	252 525	CAA GTT	GTC	AAC TTG
GAT	AAA TTT	CAT	GCT	CAT
GTC GAT	GCG GCC AAA CGC CGG TTT	TTG	GAT	ນນອ ອອນ
GTA	000 000	AAA TTT		GTC ATG
ອນອ	ე <u>ნე</u> ნენ	TAG	ACT TGA	CAG
GAT	TGG	TGC GCA TAG ACG CGT ATC	ATA GTG Z	225 552 552 225
CAT	GAC	TGC	ATA TAT	990 009
929 292	CAG	000 000	5 50	GAG
GGT	GAG	GAG AAC CTC TTG	CAC	CAA
TGT	AGC		CAG	ວ <u>ງ</u> ອນນ
ວວວ ອອອ	CGA	TCC	TAG	ATC
3106	3151	3196	3241	3286

					. •	,
	ATT	TAA	CIG		AAC	
	J D D D	909		ATT	TCA	
i	GNG	CIC	ALL	TAA	CTG	GAC
	GAT	CTA	GCA	CGI	AAG	TTC
	GAC	CTG	TTA	AAT	GAT	CTA
	GAT	CTA	SCG	ລອລ	GAT	CTA
	GNG	CTC CTA CTG	1CT	IGA	ATC	TAG
			CTG	GAC 1	CTT	GAA TAG
	CCT	CCA	TGC	ACG		TTC
176. 11	GAC	CCA CTG CCA CGG	990	ລລອ	TTA	AAT
	GGT	CCA	ACA	TGT	GCA	CGT
	CAG	GTC	CAT	GTA		TGG
	ATC	TAG	TTT	AAA	ACT	TGA
	AGC	TCG	AGA	TCT	TAA	ATT
	TAC	ATG	GTT	CAA	TGA	ACT
	3331	•	3376		3421	

43/60

Total number of bases is: 3474.

ATG AGA ATT TAC TCT TAA

3466

Total number of Dases is. 34/4.

DNA sequence composition: 845
2 OTHER;

716

Ğ:

978

933

Sequence name: NPMTNFMPH



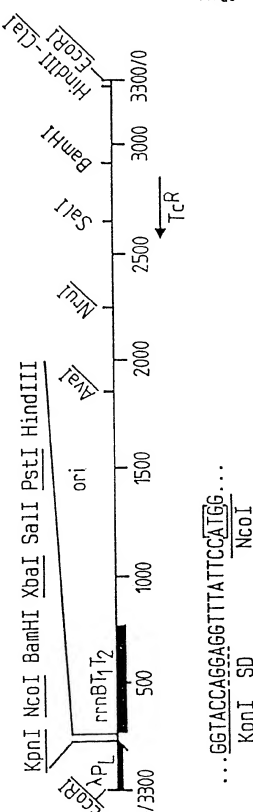


Fig. 12a

	10 A F:	~ C	<i>የ</i> ካ ደነ	45/60	d [1	F-1 e-1	F1 &	ひひ
	45 AAA TTT	AAA TTT	CTG	CTT ⁵ GAA ₉ /	AAA TTT	GCT CGA	GAT	TGG
	AAA TTT	GAT	ATA TAT	GCT	TTT AAA	CAA	ACA TGT	990 ၁၁9
	39 TGC ACG	GGT	GTG	GAC	AGG TCC	552 225	GAT	TTT AAA
	999 222	TGC	252 525	GGT	AGG TCC	GCA	CCT	GAA
	33 - - - -	ATC TAG	CTG	GAA	ACC TGG	CCT	CAG	ACA TGT
	ATG TAC	ACC	CCA	CAT	GGT	CGA	TTT AAA	AAA TTT
	27 ACA TGT	ATA	ATA TAT	CAC	AGG	AGT TCA	GAT	GAT
12b	CAA GTT	CAG	TAA ATT	GAC	ອນນ ນອອ	TAG	GAA	TCT AGA
Fig.	21 	ATA TAT	ACA TGT	ACT TGA	AAG TTC	CTC	AGA TCT	ວລອ ອອລ
	ACC TGG	AAC TTG	TTG	505 050	AAG TTC	ATC TAG	ATG TAC	AAG TTC
	15 	AAA TTT	GTG	GGA	CTG	000 000	000 000	CAG
	TCT AGA	ATA TAT	000 000	GCA	550 005	000 000	TGG	ACG
	9 	CAT	CTG	TCA	TAA	ATG	TTT AAA	AGA TCT
pIG2	ລວອ ອອລ	ATT TAA	TCT	ACA TGT	AAT TTA	TCC	CTG	ATC TAG
	3 TTC AAG	TAA	TTA	AGC TCG	AAA TTT	TAT ATA	TGG	TAA
From:	7	46	91	136	181	226	271	316

990 000

ATG

000 000

TGA

TCC

CCA

AGG

AGA TCT

AGC

TTA

AAA TTT

ATC

922

CCA

CTG

631

CAT

505 000

990 000

GAC

CAG

222 222

922 263

GGT

GAG

000 000

AAC

AGC

CGA

586

	d : F-1	r \		10 T \	46/60
	AG/	TGC	AGT TCA	ACG TGC	TTG 57
	CTC	CCA	CTC GAG	TGA	ACG
	GAA	TCC	AGG TCC	225 552	TGA
	990 229	GTC	GAA CTT	TGT ACA	ATT TAA
		222 222	AAC TTG	GTT CAA	ວວອ ອອວ
	555 222	TGT	TAA ATT	GTT CAA	GAG
	TGA	TAG	AAA TTT	TCT	ეე <u>ნ</u> ეეე
. 12b	ACC	TGG	ATC	TTA	5 25
Fig	555 ၁၁၁		_{ອວວ}	GTT CAA	ATC TAG
	GGT	909 090	CCA	TTC	CAA
	GGT	TAG	CTG	CCT	GGA
	929 260	ე <u>ე</u> ნე	GAA	000 000	GTA
	CAG TAG CGC GTC ATC GCG	ACG	AGG TCC	ACT TGA	TGA
	CAG	GAA	AGT TCA	AAG	TCC
	361 CGG GCC	406 AGT TCA	GAG	496 CGA A	CTC
	361	406	451	496	541

CAT CTT TAA TAA TGA TTC 555 555 TTT TAA TTA TGT CAA TTT AGA TCT CTCATG CTC AAA TTT TAC 255 522 TTC TAT ATG CGT TTG AAT TTT AAA

CAT GTA	AGA TCT	S TTG AGA TCC TTT TTT TCT	AGC 09/	GAA CTT	TCT AGA	ACC TGG
TCT	GTC	TTT AAA	ACC TGG	TTT TCC AAA AGG	CCT	AGC TCG
raa att	AGC	TTT AAA	GCT ACC CGA TGG	TTT AAA	GAT ACC AAA TAC TGT CCT CTA TGG TTT ATG ACA GGA	CTC TGT AGC GAG ACA TCG
TTT TGA AAA ACT	CTG	TCC AGG	ACC ACC	TCT AGA	TAC	CTC
TTT AAA	CCA	AGA TCT	ACC	GCT ACC AAC CGA TGG TTG	AAA TTT	GAA CTT
CCT	GTT CAA	TTG	AAA TTT	ACC TGG	ACC TGG	CAA GTT
T &	GTT TTC CAA AAG	rTC AA(GCA AAC AAA CGT TTG TTT	GCT	GAT	CTT
GAA GACTT CT	GTT CAA	AGG ATC 1 TCC TAG A	AAC TTG	AGA TCT	AGC GCA TCG CGT	CCA
Fig. 1 GGT CCA	TGA ACT	AGG		TCA	AGC	CCA
CTA GAT	ACG	CAA GTT	CTT GAA	GGA	CAG	AGG TCC
GAT	TTA	GAT	CTG	990 009	CAG	GTT
AAG TTC	၁၁၁	AAA TTT	CTG	TTT AAA	CTT	GTA
TAA	AAT TTA	AGA TCT	AAT TTA	TTG	TGG	990 000
TAA ATT	CAA GTT	CGT	CGT	GGT	AAC	GTA
CAA GTT	GAC	555 555	ວຽວ ອວອ	GGT	GGT	1036 AGT TCA
992	811	856	901	946	991	1036

AGA TCT TAG ATC

AGG TCC TTA AAT

AAC TTG TCT AGA

CGG GCC GTA GTA

GGT CCA CTG GAC

CAG GTC CGC GCG

CGG GCC AAA TTT

AAG TTC GGG CCC

GGT CCA AGG TCC

TCC AGG TCC AGG

GTA CAT GCT CGA

GGC CCG CAC GTG

GGA

GAG CTC

292 929

1351

CAG

GGA

AAA TTT

1306

t)	
Con	
12b (
Fig. 12	

TGC	ATA TAT	GTG CAC ₂	ATA [©] TAT	GAG		
			GAG	AGG TCC		
922 299		222 999	ACT TGA	CGA		
AGT TCA	CTC	222 222	CGA	TCC		
ACC	GGA	AAC TTG	CAC	GCT		
GTT	GTT CAA	CTG	CTA GAT	CAC GCT GTG CGA		
	ນນອ ອອນ	ວວວ ອອອ	GAC	505 050		
	TAC	GTC	AAC TTG	AAG TTC		
	TCT AGA	090 909	ე <u>ნე</u>	AGA TCT		
	GTG	GCA	GGA	TTG		
505 050	GTC	522 255	CTT	GCA		
CCT	TAA ATT	TAA	CAG	TGA		
ATA TAT	CGA	GGA	990	ວອວ ອວອ		
TAC	TGG	ACC	ACA TGT	ACA		
99 0 ၁၁9	CAG		CAC	CCT		
1081	1126	1171	1216	1261		
SUBSTITUTE SHE						
	CCT	1081 GCC TAC ATA CCT CGC TCT GCT AAT CCT GTT ACC AGT GGC TGC CGG ATG TAT GGA GCG AGA CGA TTA GGA CAA TGG TCA CCG ACG ACG ATG TAT GGA GTC GTG TCT TAC CGG GTT GGA CTC AAG ACG GTC ACC ACC ACC ATT CAG CAC AGA ATG GCC CAA CCT GAG TTC TGC	1081 GCC TAC ATA CCT CGC TCT GCT AAT CCT GTT ACC AGT GGC TGC CGG ATG TAT GGA GGG TTA GGA CAA TGG TCA CCG ACG ACG ATG TAT CAG CGC ACG TTA GGA CAC TGC ACG ACG GTC ACG CGT ATT CAG CAC AGA ATG GCC CAA CCT GAG TTC TGC CAA CCT GAG TTC TGC CAA CCT GAG TTC TGC CAA TGC CCT ATT CCG CCT CGC CTC CAG CCC CAA CCC CAA CCC CCC AAG	1081 GCC TAC ATA CCT CGC TCT GCT AAT CCT GTT ACC AGT GGC TGC CGG ATG TTA GGA CGA TTA GGA CAA TGG TCA CCG ACG ACG ACG ATG TTA GGA CAA TGG TCA CCG ACG ACG ATG TAA GTC GTG TCT TAC CGG GTT GGA CTC AAG ACG TC ATT CAG CAC AGA ATG GCC CAA CCT GAG TTC TGC CAA TTC CAG CAC GCG GTC GGG CTG AAC GGG GGG TTC TAC CAA TGG CCT ATT CGG CGT CGC CAG CCC GAC TTG CCC CAA CCC GAC TTG CCC CAG TTG CTG GAT TGA CTC GAA CTC CGA ACT GAG CTC TGA CTC CAG TTG CTG CTG TGA CTC CAG CTC TGA CTC		

	49/60							
	GTG	929 292	CAT GTA	TAC	CGA			
(2 120 (2011 6)	TTT AAA	CNA GTT	TCA	TAT ATA	GAC			
		CAG	TGC	ე <u>ე</u> ნე	AAC TTG			
	TCG ATT AGC TAA	525 252	TTT AAA	TAA ATT	ວ <u>ອ</u> ອ ອວວ			
	252 525	AAA TTT	550 225	GGA	CAG			
	TGA	GAA	CTG	TGT ACA	299 922			
	ACT	ATG	TTG	TTC	TCG			
	CTG	CCT	CTT GAA	TGA	909 090			
	CCT	GAG CTC	ອນນ	999	TAC			
	CCA GGT	292 929	CCT	ATC TAG	TGA			
	TCG AGC	222 255	GTT CAA	GTT CAA	AGC TCG			
	GTT	AGG	ACG	TGC	GTG			
	CGG GTT GCC CAA	GTC	TTT AAA	TCC TGC AGG ACG	TGA GTG ACT CAC			
	TGT ACA	CTC	CTT GAA	CTT GAA	CTT			
	TCC AGG	ATG TAC	GGC CTT CCG GAA	GTT CAA	505 050			
	1396 TCC TGT AGG ACA	1441 ATG TAC	1486	1531 GTT CAA	1576 CGC CTT GCG GAA			

49/60

SUBSTITUTE SHEET									
1621	1666	1711	1756	1801	1846	1891			
ນອ ນ ອນອ	ວ <u>ອ</u> ວ ອວອ	TTG	TTC	999 222	GCA	522 255			
CAG	TTT AAA	TTG	GCT CGA		999 ၁၁၁	TGC			
CGA	CCA	CTC	505 050	GCC AGC CGG TCG	GTG	TGG			
GTC	GAC	AGG	GTA	CTA	990 009	AGA			
AGT TCA	TTT AAA	TCG	TCG	ອອວ ວວອ	AGG	TGG			
GAG	ACG	CAG	GTG	323 333	ACC	555 555			
Fig. 1 CGA GCT	AAA TTT	ACG TGC	ATT TAA	TCC AGG	CAA	ACG TGC			
GGA CCT	CAC	TTT	CAT		ອນອ	CGG ACG CGA GCC TGC GCT			
Fig. 12b (Con't) CGA GGA AGC GCT CCT TCG	CAC GGA A	TGC	TCT AGA	ACG	TGC	TGG			
GGA	AAC FTG	TGC AGC AGC AGT ACG TCG TCG TCA	CAT TCT GCT AAC GTA AGA CGA TTG	TCA ACG ACA AGT TGC TGT	CGC TGC CCG AGA TGC GCC GCG GCG ACG GGC TCT ACG CGG CGC	TGG ATA TGT TCT GCC AAG ACC TAT ACA AGA CGG TTC			
AGA TCT	CGA AGA (GCT TCT (AGC	AAC TTG	GGA	AGA TCT	TGT			
ე ეე	AGA TCT	AGT TCA	CAG	GCA (CGT)	TGC	TCT			
CTG	CCA GGT))))	CAG TAA (GTC ATT (CGA	9 90	990 009			
ACT TGA	TTC	TTC	9 22	CGA TCA GCT AGT	ວ <u>ອ</u> ວ ອວອ	AAG TTC			
CTG ACT TCC GAC TGA AGG	ATG	ACG TGC	A GGC AAC ^S r CCG TTG	TGC	TGC	GGT			

CAA GTT	CAT GTA	55 555	1/60 りりり ンンン	CAG	TCC AGG	CAT
	TC	ອນອ ນອນ	CAA GTT	GAT	CGA	CAG
TGG	5 22	CAA GTT	TGC	GAC CTG	GAG CGA	GGA CAG CCT GTC
GAT TGG CTC CTA ACC GAG	7 900 990 1 099 009	ACG TGC	CCA	CGT	929 292	CCT
ATT	990	ე <u>ეე</u> ე	AAT TTA	909 090	AGC	CTG
AGA TCT	GGT	TGC ACC GCG ACG CAA ACG TGG CGC TGC GTT	GCC TAC AAT CCA TGC CAA CCC CGG ATG TTA GGT ACG GTT GGG	AAT CGC C TTA GCG 6	AAG TTC	TAC
GCA CGT	CGA	TGC	550 005	ATA	GGT	ATC TAG
Fig. 12b (Con't) TTC TCC GCA AGA AAG AGG CGT TCT	CGT TAG CGA GGT GCA ATC GCT CCA	CCA	500 055	ອນນ	GCT GGT AAG AGC CGC CGA CCA TTC TCG GCG	GTC ATC TAC CTG
TTC AAG	CGT	CCG GCT GGC CGA	500 500 055 055	ອນນ	TAG	CTG ATG GTC GAC TAC CAG
CAG	ATC TAG	ე <u>ე</u> ნე	TAG ATC	CGA	AGT TCA	ATG
TCA	TGA	922 299	GTA	ອນອ ນອນ	CGA	
CAT	TGG	GGT CCA	AAG TTC	GCT	GAT	TCC
ນ ອນອ	GAG	CGA	GAC	TGT	AGT	CTG
LTT AAA	TTG	GGT CCA	GCA	CCA	TCC	AAG TTC
TGG	TTC	TCA AGT	GAG	GTT CAA	000 000	TTG
1936	1981	2026	2071	2116	2161	2206
					_	

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	AAT	TTA	CAA	52/6 LLS	CTG	GAC	TTG	AAC	GAT	CTA
	AAG			GTC	CCC	೮೦೦	299	ອວວ	229	990
	GAG			929	AAT	TTA	GAA	CTT	CAG	GTC
	AGC	TCG		CLL	GGC GAT	CTA	GAC	CTG	CGA	GCT
	GGA	CCL	CGC	929	299	ອນນ			AAG	TTC
	225	990	CGT	GCA	229	SSO	ACC	CCC IGG ICA	CGC	909
con.t)	225	SSS	TCG	; AGC GCA	CAT	GTA	999	ວລວ	TAC	ATG
rig. IZD (Con.t.	GAT	CTA	229	ອອວ	CGC	ອວອ	CGC	ອວວ	GAA	CTT
F18	၁၁၁	999		GGT	CCC	ອວວ		CCA	TCC	AGG
	CAT	GTA		GTA	GTC	CAG	TTT	AAA	GAT	CTA
	999			ອວວ		೮೦೮	ACG	TGC	CAA	GTT
	၁၅၁	909	GAA	CTT	CAG	GIC	GAA	CTT	GTG	CAC
	CTG CAA	GTT	CCC	ນນນ	GCC CAG	ອອວ	CCC		GAG GGC	SSS
	CIG	GAC	AAT	ľTA	GTA	CAT	CIC		GAG	CIC
	SGC	ອນນ	CAT	GTA	GAC	CTG	CTT	GAA	AGC	TCG
	2251 GGC (2296	GTA	2341 GAC GTA		2386		2431 AGC (

CCA	GAC	GNA CTT	CTC	GTT CAA	GGA	ACC TGG
GAC	GAA	299 922	ACG	TAG	CAA GTT	CAT
AAT TTA	AAA TTT	CCA	CGG TCG ACG GCC AGC TGC	GCC CAG TAG CGG GTC ATC	TGG TGC ATG ACC ACG TAC	CAC
GAA	SAT		225 552	CAG	TGC	CAC GGG GCC TGC GTG CCC CGG ACG
990 009	CAT (GTA	ე <u>ე</u> ნე	CAT	ອອວ ວວອ	TGG	9 90
	TTG	909 099 990 090 900 009	GGG CAT	GCA	GAA	ວວວ
Fig. 12b (con't) GCG GTC CTC CGC CAG GAG	GAG	GAT AGT CAT CTA TCA GTA	GGC TCT CAA CCG AGA GTT	GAA	AAG TTC	CAC
Fig. 12 GCG CGC	TAC ATG	AGT TCA	TCT	TAG	2 525 525 2 525 525	922 999 299 222
ANA TTT	TCC	GAT		CAT	505 050	999 222
ນ <u>ອ</u> ນ	CTG	GAC	GAA	CTG	525 252	TCC
CCA	CAC	922 299	GTT CAA	CTC	CAC	CAG
GCT	ລວອ	TGC	TGG	CGA	GAG	CAA
ອນອ	TGC	AAG TTC	GAC	ATG TAC	GTT	550
CGT	505 050	CAT	GCT	CTT	950 205	922 299
CAT	GAG	AGT TCA	GGA	TCC	GAG	GAT
2476	2521	2566	2611	2656	2701	2746

Fig. 12b (Con't)

CGG TTT CGC CAG TTG CAT CAA CGC	GTA GTT GCT GTC CGA CAG
G TTT CGC G CAT CAA	GCT
G TTT G CAT	GTA GCT CGA
ט טנ	
CGC	AAC GAT CTA
3GC AAA	TTT
CGC	ATC ACT TGA
TGG ACC GCA	GTG ACT
GAC CTG TGC	CCC ACG GCC ATA CGG TAT
CAG GTC GGG	990 009 000
	TTG
	CTC
	AGG
	GTC TAG ATC
TCC AGG	CCT ATA TAT
2971	
	TCC AAG TAG CGA AGC GAG CAG GAC TGG GCG GCG AGG TTC ATC GCT TCG CTC GTC CTG ACC CGC CGC

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't)
Con.
12b (
Fig.

			5	5/60
CAA	GAT	GCA	AAG TTC	
AAC TTG	GAC	TTA	GAT	
CAT	GAT	252 525	GAT	
225 252	GAG	ACT	ATC TAG	
TAC	550 225	CTG	CTT GAA	
CAG	GGT CCA	TGC	AAG TTC	
225 552	GAC	ນວອ	TTA	
550	GGT	ACA TGT	GCA	
GAG	CAG	CAT	ACC	
CAA GTT	ATC TAG	TTT AAA	ACT TGA	A T
255 522	AGC TCG	AGA TCT	TAA	AGA A TCT T
ATC	TAC	GTT CAA	TGA	ATG
GAC GAT CTG CTA	990 ၁၁9	ATT TAA	TAA CTG ATT GAC	TCA AAC ATG AGT TTG TAC
GAC	TAT ATA	CGC ATT GCG TAA	TAA	
ATG	GCC TAT C		ATT TAA	CTG
3106 ATG TAC	3151	3196 GAG CTC	3241	3286

681 G; 936 887 A; 3301. Total number of bases is: DNA sequence composition:

Sequence name: NIPS0039.

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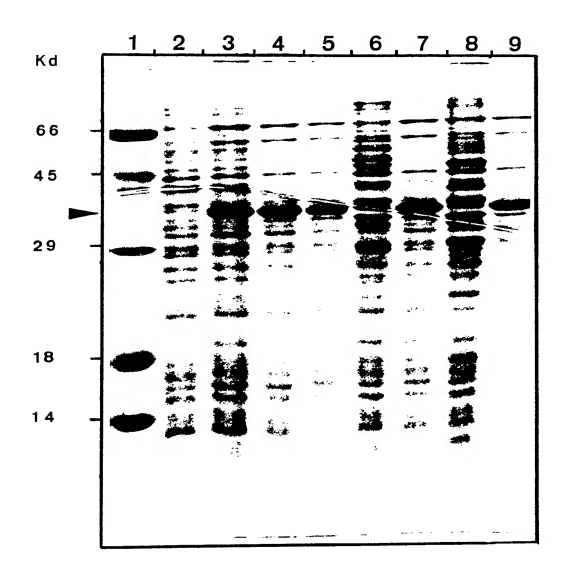


fig. 14a

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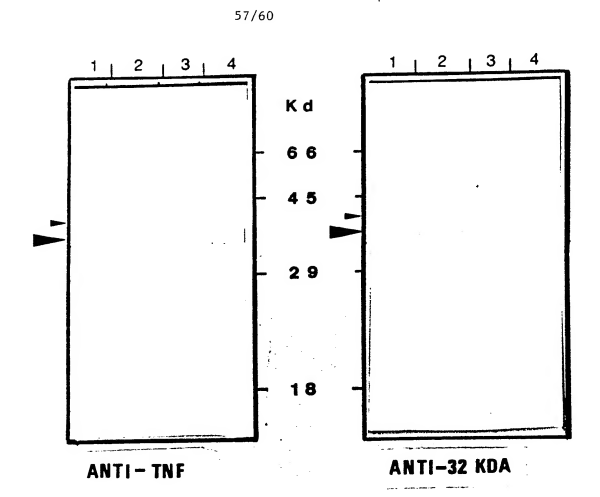
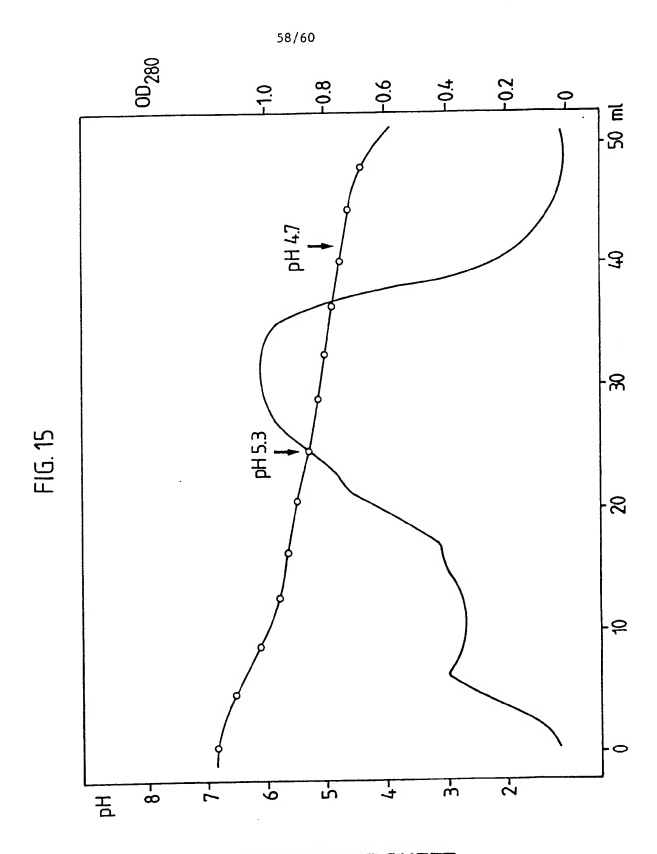
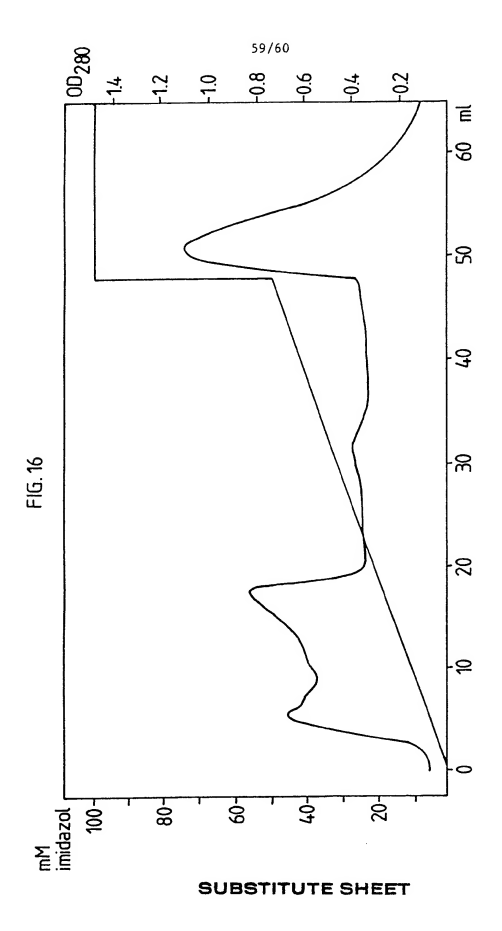


fig.14b

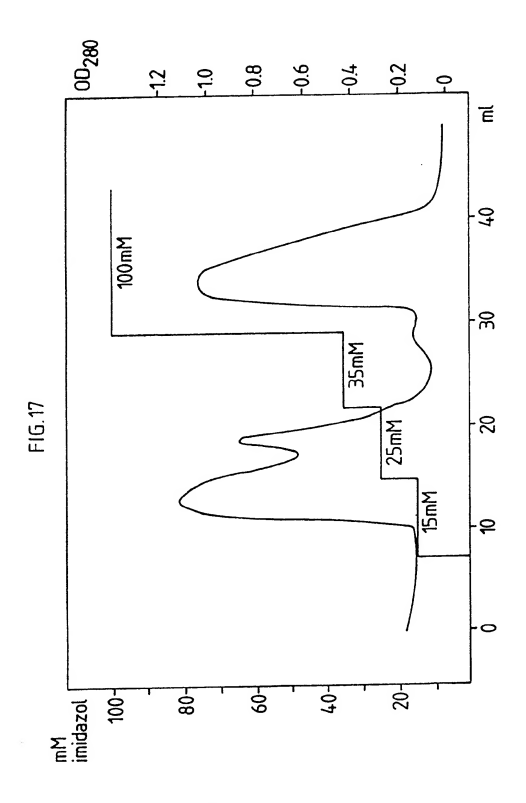
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INTERNATIONAL SEARCH REPORT

International Application No PCT/EP 90/01593

		International Papers	/ 11
I. CLASSI	FICATION OF SUBJECT MATTER (if several classifi	cation symbols apply, indicate all) 6	
According	to International Patent Classification (IPC) or to both Natio	onal Classification and IPC	
. 5	C 07 K 13/00, A 61 K 39/04	l, C 12 N 15/31, G	01 N 33/569
IPC:	C 12 Q 1/68		
II. FIELDS	SEARCHED		
	Minimum Document	tation Searched 7	
Classification	n Svetem 1	Classification Symbols	
Liassification	1 System (
IPC ⁵	C 07 K, C 12 N, A 61 K	C, G 01 N, C 12 Q	
	Documentation Searched other the to the Extent that such Documents	nan Minimum Documentation are included in the Fields Searched ⁸	
III. DOCUI	MENTS CONSIDERED TO BE RELEVANT		Relevant to Claim No. 13
ategory •	Citation of Document, 11 with Indication, where appr	opriate, of the relevant passages 12	Refevent to Claim No.
j			
x	Chemical Abstracts, volum 1983, (Columbus, Ohio H. Tasaka et al.: "Pu antigenic specificity (Yoneda and Fukui) fr tuberculosis and myco intracellulare", see page 413, abstract & Hiroshima J. Med. St 1-8 (Eng). cited in the application	o, US), drification and of alpha protein com mycobacterium bacterium et 86251m aci. 1983, 32(1),	1-9,40,41
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X	Journal of Clinical Micr 25, no. 7, July 1987, M.L. Cohen et al.: "proteins of mycobacte in escherichia coli a recombinant genes and development of diagno page 1176 see the whole documen cited in the application	Expression of rium tuberculosis nd potential of proteins for stic reagents",	10-22,25-33,35-39,43,44
4.6	l cottogories of cited documents: 10	"T" later document published after t	he international filing date
"A" docu cons "E" earli filing "L" docu	I categories of cited documents: 10 ument defining the general state of the art which is not sidered to be of particular relevance er document but published on or after the international g date ument which may throw doubts on priority claim(s) or call its cited to establish the publication date of another tion or other special reason (as specified)	or priority date and not in conflicted to understand the principli invention "X" document of particular relevan cannot be considered novel of involve an inventive step "Y" document of particular relevan	ce; the claimed invention ce; the claimed invention cannot be considered to ce; the claimed invention an inventive step when the
"O" doci	ument referring to an oral disclosure, use, exhibition or or means ument published prior to the international filing date but	document is combined with one ments, such combination being in the art.	phylous to a person skilled
later	r than the priority date claimed	"&" document member of the same	hereur sermit
IV. CERT	IFICATION		
	Actual Completion of the International Search	Date of Mailing of this International S	
	20th December 1990	23 JAN	וַבְקוֹ
Internation	at Searching Authority	Signature of Authorized Officer	HW XX
	EUROPEAN PATENT OFFICE	Mme N. KUIPER -	1

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х	BE, A, 905582 (INSTITUT PASTEUR) 9 April 1987 see pages 10,11; claims	35-39,45
х	Journal of Bacteriology, volume 170, no. 9, September 1988, Am. S∞. for Microbiology, K. Matsuo et al.: "Cloning and expression of the mycobacterium bovis BCG gene for extracellular antigen pages 3847-3854	10-22,40,41 43
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Y	EP, A, 0288306 (McFADDEN) 26 October 1988 see page 7, column 12, lines 2-18	23,24,32,3 42
A	<pre>Int. Archs Allergy appl. Immun, volume 81, 1986, S. Karger AG (Basel, CH), H.G. Wiker et al.: " MPB59, a widely cross-reacting protein of mycobacterium bovis BCG", page 307</pre>	-
A	Microbial Pathogenesis, volume 2, 1987, Academic Press Inc. (London GB) Ltd., J. De Bruyn et al.: "Purification characterization and identification of a 32 kDa protein antigen of mycobacterium bovis BCG", pages 351- 366	
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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

EP 9001593

SA 40401

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 16/01/91

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

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BE-A- 905582	09-04-87	None		
EP-A- 0288306	26-10-88	AU-A- EP-A- WO-A-	0356450	07-03-90

For more details about this annex: see Official Journal of the European Patent Office, No. 12/82